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Date: October 27, 2000

Docket No.: 20-4764P



Assistant Commissioner for Patents Washington, DC 20231

Sir:

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As authorized by the inventor(s), transmitted herewith for filing is a Continuation-in-part application of Appl. No. 09/302,357 filed on April 30, 1999.

Inventor(s): NAKAJIMA, Hiroki and NAGASAWA, Akitsu

For: METHOD FOR GIVING RESISTANCE TO WEED CONTROL COMPOUNDS TO PLANTS

Enclosed are:

⊠ A pages	A	specification	consisting	of	three	hundred	thirteen	(313)
		irtoon (12)						(010)

- \boxtimes Thirteen (13) sheet(s) of formal drawings
- Executed Declaration (\boxtimes Original \square Photocopy) M
 - Preliminary Amendment
- An assignment of the invention to Sumitomo Chemical Co., \boxtimes
- Certified copy of
- Priority of application No(s). 120553/1998 filed in JAPAN on April 30, 1998 is claimed under 35 U.S.C. \$ 119. See attached copy of the Letter claiming priority filed in the prior application on _____.

	\boxtimes	An Information Disclosure Statement and PTO-1449 form(s) are attached hereto for the Examiner's consideration.						
		A statement (Original Photocopy) to establish small entity status under 37 C.F.R. § 1.9 and 37 C.F.R. § 1.27.						
		Amend to		cation h	y inse	rting bef	ore the	first
		This application is a continuation-in-part of Application No. 09/302,357 filed on April 30, 1999, the entire contents of which are hereby incorporated by reference						
a Kin Cin	Amend the specification by inserting befo sentence						ore the	first
		This application is a continuation-in-part of PC international application No. PCT/_ that has a international filing date of that designated th United States, the entire contents of which are hereb incorporated by reference						
ロチウンティウ		An extension of time for () month(s) until has been submitted in parent Application No. 09/302, $\overline{357}$ in order to establish copendency with the present application.						
Ē		Other:						
heal	The filing fee has been calculated as shown below:							
T.	The filing fee has been calculated as shown below: LARGE ENTITY SMALL BASIC FEE \$710.00 \$355							
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			LARGE ENTITY		SMALL ENTITY	
BASIC F		FEE	\$710.00		\$355.00	
	NUMBER FILED	NUMBER EXTRA	RATE	FEE	RATE	FEE
TOTAL CLAIMS	79-20 =	59	x 18 = \$1062.0		x 9 =	\$0.00
INDEPENDENT CLAIMS	3-3 =	0	x 80 =	\$0.00	x 40 =	\$0.00
MULT _YES	+ \$270.00		+ \$135.00			
TOTAL			\$2,0	142.00	\$0.	00

GMM/rem

20-4764P Attachments

- A check in the amount of \$2,082.00 to cover the filing fee and recording fee (if applicable) is enclosed.
- Please charge Deposit Account No. 02-2448 in the amount of \$0.00. A triplicate copy of this transmittal form is enclosed.
- Please send correspondence to:

BIRCH, STEWART, KOLASCH & BIRCH, LLP or Customer No. 2292 P.O. Box 747

Falls Church, VA 22040-0747 Telephone: (703) 205-8000

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. § 1.16 or under 37 C.F.R. § 1.17; particularly, extension of time fees.

Respectfully submitted,

BIRCH, STEWART, KOLASCH & BIRCH, LLP

By Gerald M. Murphy, Jr., #28,977

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(Rev. 09/29/00)

IN THE U.S. PATENT AND TRADEMARK OFFICE

Applicant:

NAKAJIMA, Hiroki et al.

Appl. No.:

NEW

Group:

Filed:

October 27, 2000

Examiner:

For:

METHOD FOR GIVING RESISTANCE TO WEED

CONTROL COMPOUNDS TO PLANTS

PRELIMINARY AMENDMENT

Assistant Commissioner for Patents Washington, DC 20231

October 27, 2000

Sir:

The following preliminary amendments and remarks are respectfully submitted in connection with the above-identified application.

AMENDMENTS

IN THE CLAIMS:

Please amend the claims as follows:

CLAIM 24: Line 1, change "claim 22 or 23" to

--Claim 22-

CLAIM 25: Line 1, change "claim 22 or 23" to

--Claim 22--

CLAIM 26: Line1, change "claim 22 or 23" to

--Claim 22-

 ${\bf CLAIM}$ 27: line 1, change "claim 22 or 23" to

--Claim 22--

REMARKS

The claims have been amended to delete improper multiple dependencies.

Entry of the above amendments is earnestly solicited. An early and favorable first action on the merits is earnestly solicited.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. § 1.16 or under 37 C.F.R. § 1.17; particularly, extension of time fees.

Respectfully submitted,

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(Rev. 04/19/2000)

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METHOD FOR GIVING RESISTANCE TO WEED CONTROL COMPOUNDS TO PLANTS

BACKGROUND OF THE INVENTION

FIELED OF THE INVENTION

The present invention relates to a method for giving resistance to weed control compounds to plants.

DISCLOSURE OF THE RELATED ART

Weed control is very important work for improving yields and quality of cultivated plants. For this purpose, weed control compounds such as herbicides are mainly used. However, for using weed control compounds, it is not always easy to distinguish cultivated plants from weeds of allied species to selectively control only weeds. Then, production of plants having resistance to weed control compounds (hereinafter referred to as weed control compound-resistance) has been attempted and some resistant plants have been put to practical use.

Recently, gene engineering techniques have been utilized for producing plants having weed control compound-resistance. As such a technique, for example, Hinchee, M.A.W. et al. disclose a method for producing a plant having resistance to a herbicide, glyphosate, wherein 5-enolpyruvylshikimate-3-phosphate synthase (EPSPS) gene

which is a target enzyme of glyphosate is mutagenized so that an affinity for glyphosate is reduced, and the gene is introduced into a plant [Hinchee, M.A.W. et al., BIO/TECHNOLOGY, 6: p 915 (1988)].

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OBJECTS OF THE INVENTION

Varieties of known methods for giving weed control compound-resistance to plants are not necessarily sufficient and it has been desired to develop further various kinds of methods for giving weed control compound-resistance to plants.

The main object of the present invention is to provide a new kind of a method for giving weed control compound-resistance to plants.

This object as well as other objects and advantages of the present invention will become apparent to those skilled in the art from the following description with reference to the accompanying drawings.

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BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 is the restriction map of plasmid pETBCH. bchH is magnesium chelatase protoporphyrin IX binding subunit gene of a photosynthetic bacterium *Rhodobacter sphaeroides*. T7 pro represents the promoter sequence of T7 phage, and T7 ter represents the terminator sequence of T7

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phage. $\mbox{Amp}^{\rm r}$ is an ampicillin resistant gene, $\mbox{lacI}^{\rm q}$ is a repressor protein gene of a lactose operon, and ori is the replication origin.

Fig. 2 is the restriction map of plasmid pACYCSP. PPO is protoporphyrinogen IX oxidase gene of soybean and lac pro represents the promoter sequence of a lactose operon. Cmr is a chloramphenical resistant gene and ori is the replication origin.

Fig. 3 is the restriction map of plasmid pTVBCH. bchH is magnesium chelatase protoporphyrin IX binding subunit gene of the photosynthetic bacterium Rhodobacter sphaeroides. lac pro represents the promoter sequence of a lactose operon. Ampr is an ampicillin resistant gene and ori is the replication origin.

Fig. 4 is the restriction map of plasmid pBIBCH. is magnesium chelatase protoporphyrin IX binding subunit gene of the photosynthetic bacterium Rhodobacter sphaeroides. NP is the promoter sequence of a nopaline synthase gene, NT is the terminator sequence of the nopaline synthase gene, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII represents a kanamycin resistant gene, and RB and LB represent right and left border sequences of T-DNA, respectively.

Fig. 5 is the restriction map of plasmid pNO. is the promoter sequence of a nopaline synthase gene, NT is

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the terminator sequence of the nopaline synthase gene, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII represents a kanamycin resistant gene, and RB and LB represent right and left border sequences of T-DNA, respectively.

Fig. 6 is the restriction map of plasmid pTCHLH. TCHLH is protoporphyrin IX binding subunit gene of tobacco magnesium chelatase whose chloroplast transit signal has been deleted. lac pro represents the promoter sequence of a lactose operon. Amp' is an ampicillin resistant gene, Km' is a kanamycin resistant gene and ori is the replication origin.

Fig. 7 is the restriction map of plasmid pBITCHLH. TCHLH is protoporphyrin IX binding subunit gene of tobacco magnesium chelatase whose chloroplast transit signal has been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of the nopaline synthase and 35S is the 35S promoter of cauliflower mosaic virus. NPTII represents a kanamycin resistant gene, and RB and LB represent right and left border sequences of T-DNA, respectively.

Fig. 8 is the restriction map of plasmid pTVGMP. GMP is soybean protoporphyrinogen IX oxidase gene whose chloroplast transit signal and FAD binding sequence have been deleted. lac pro represents the promoter sequence of

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a lactose operon. Amp' represents an ampicillin resistant gene and ori is the replication origin.

Fig. 9 is the restriction map of plasmid pBIGMP. GMP is soybean protoporphyrinogen oxidase gene whose chloroplast transit signal and FAD binding sequence have been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 10 is the restriction map of plasmid pTVCRP. CRP is protoporphyrinogen oxidase gene of *Chlamydomonas* reinhardtii whose chloroplast transit signal and FAD binding sequence have been deleted. lac pro represents the promoter sequence of a lactose operon. Amp' is an ampicillin resistant gene and ori is the replication origin.

Fig. 11 is the restriction map of plasmid, pBICRP.

CRP is protoporphyrinogen oxidase gene of *Chlamydomonas*reinhardtii whose chloroplast transit signal and FAD

binding sequence have been deleted. NP is the promoter

sequence of a nopaline synthase, NT is the terminator

sequence of a nopaline synthase, and 35S is the 35S

promoter of cauliflower mosaic virus. NPTII is a kanamycin

resistant gene, and RB and LB are the right and left border

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sequences of T-DNA, respectively.

Fig. 12 is the restriction map of plasmid pTVHVF1. HVF is barley ferrochelatase gene whose signal sequence has been deleted. lac pro represents the promoter sequence of a lactose operon. Amp' represents an ampicillin resistant gene and ori is the replication origin.

Fig. 13 is the restriction map of plasmid pBIHVF. HVF is barley ferrochelatase gene whose signal sequence has been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 14 is the restriction map of plasmid pTVCSF. CSF is cucumber ferrochelatase gene whose signal sequence has been deleted. lac pro represents the promoter sequence of a lactose operon. Amp^c is an ampicillin resistant gene, and ori is the replication origin.

Fig. 15 is the restriction map of plasmid pBICSF.

CSF is cucumber ferrochelatase gene whose signal sequence has been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant

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gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 16 is the restriction map of plasmid pHEMF. HEMF is coproporphyrinogen III oxidase gene (hemF) of Escherichia coli. lac pro is the promoter sequence of a lactose operon. Amp^c is an ampicillin resistant gene, and ori is the replication origin.

Fig. 17 is the restriction map of plasmid pBIHEMF. HEMF is coproporphyrinogen III oxidase gene (hemF) of Escherichia coli. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 18 is the restriction map of plasmid pBIHASYS8. HASYS8 is a gene encoding MG(HASYS)₈ protein. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 19 is the restriction map of plasmid pBIRASSL8. RASSL8 is $MG(RASSL)_g$ protein. NP is the promoter sequence of a nopaline synthase, NT is the

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terminator sequence of a nopaline synthase, and 35s is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 20 is the restriction map of plasmid pNATP. PPO(A220V) is a PPO gene having a herbicidal compound-resistant mutation (A220V). NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 21 is the restriction map of plasmid pBIAPTCH. PPO(A220V) is a PPO gene having a herbicidal compound-resistant mutation (A220V) and TCHLH is a tabacco magnesium chelatase subunit gene whose chloroplast transit singal has been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 22 is the restriction map of plasmid pCRATF.

ATF is a chloroplast-localized type ferrochelatase gene of
Arabidopsis thaliana. lac pro represents the promoter

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sequence of a lactose operon. Amp $^{\rm r}$ is an ampicillin resistant gene, ${\rm Km}^{\rm r}$ is kanamycin resistant gene and ori is the replication origin.

Fig. 23 is the restriction map of plasmid pBIATF. ATF is a chloroplast-localized type ferrochelatse gene of Arabidopsis thaliana. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 24 is the restriction map of plasmid pBIAPATF. PPO(A220V) is PPO gene hvaing a herbicidal compound-resistant mutation (A220V). ATF is a chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 25 is the restriction map of plasmid pCRSCPOX. SCPOX is soybean coproporphyrinogen III oxidase gene and lac pro represents the promoter sequence of a lactose operon. Amp $^{\rm r}$ is an ampicillin resistant gene, Km $^{\rm r}$ is a kanamycin resistant gene and ori is the replication

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origin.

Fig. 26 is the restriction map of plasmid pBISCPOX. SCPOX is soybean coproporphyrinogen III oxidase gene. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 27 is the restriction map of plasmid pBIAPSCP. PPO(A220V) is PPO gene having a herbicidal compound-resistant mutation (A220V). SCPOX is soybean coproporphyrinogen III oxidase gene. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 28 is the restriction map of plasmid pCREPSPS. CTP-EPSPS is a variant gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. lac pro represents the promoter sequence of a lactose operon. Ampr is an ampicillin resistant gene, Kmr is kanamycin resistant gene and ori is the replication origin.

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Fig. 29 is the restriction map of plasmid pNG01. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, GUS is β -glucuronidase gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 30 is the restriction map of plasmid pNG04. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, GUS is β -glucuronidase gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 31 is the restriction map of plasmid pNT35S. NT is the terminator sequence of a nopaline synthase, 35S is the 35S promoter of cauliflower mosaic virus, and lac pro is the promoter sequence of a lactose operon. Amp^r is an ampicillin resistant gene and ori is the replication origin.

Fig. 32 is the restriction map of plasmid pCENS. CTP-EPSPS is a variant gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. NT is the terminator

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sequence of a nopaline synthase, 35S is the 35S promoter of cauliflower mosaic virus, and lac pro is the promoter sequence of a lactose operon. Ampr is an ampicillin resistant gene, ori is the replication origin.

Fig. 33 is the restriction map of plasmid pCENSK. CTP-EPSPS is a variant gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. NT is the terminator sequence of a nopaline synthase, 35S is the 35S promoter of cauliflower mosaic virus, and lac pro is the promoter sequence of a lactose operon. Amp^r is an ampicillin resistant gene, ori is the replication origin.

Fig. 34 is the restriction map of plasmid pBICE. CTP-EPSPS is a variant gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and PB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 35 is the restriction map of plasmid pBICETCH. CTP-EPSPS is a variant gene in which EPSPS gene

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derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. TCHLH is protoporphyrin IX binding subunit gene of tabacco magnesium chelatase whose chloroplast transit singal has been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 36 is the restriction map of plasmid pBIGMP. GMP is soybean PPO gene whose chloroplast transit signal and FAD binding sequence have been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 37 is the restriction map of plasmid pBICEGMP. CTP-EPSPS is a chimera gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. GMP is soybean PPO gene whose chloroplast transit signal and FAD binding gene have been deleted. NP is the promoter sequence of a

nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 38 is the restriction map of plasmid pBICRP. CRP is PPO gene of *Chlamydomonas reinhardtii* whose chloroplast transit signal and FAD binding sequence have been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 39 is the restriction map of plasmid pBICECRP. CTP-EPSPS is a variant gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. CRP is PPO gene of Chlamydomonas reinhardtii whose chloroplast transit signal and FAD binding sequence have been deleted. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and

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left border sequences of T-DNA, respectively.

Fig. 40 is the restriction map of plasmid pBICEATF. CTP-EPSPS is a variant gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS originated from petunia. ATF is chloroplast-localized type ferrochelatase gene of Arobidopsis thliana. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 41 is the restriction map of plasmid pBISCPOX. SCPOX is soybean coproporphyrinogen III oxydase gene. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

Fig. 42 is the restriction map of plasmid pBICESCPOX. CTP-EPSPS is a variant gene in which EPSPS gene derived from Agrobacterium is ligated to the downstream of a nucleotide sequence encoding a chloroplast transit peptide of EPSPS derived from petunia. SCPOX is

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soybean coproporphyrinogen III oxydase gene. NP is the promoter sequence of a nopaline synthase, NT is the terminator sequence of a nopaline synthase, and 35S is the 35S promoter of cauliflower mosaic virus. NPTII is a kanamycin resistant gene, and RB and LB are the right and left border sequences of T-DNA, respectively.

SUMMARY OF THE INVENITON

Under these circumstances, the present inventors have studied intensively so as to develop a new kind of a method for giving weed control compound-resistance to plants. As a result, it has been found that weed control compound-resistance can be given to plants by allowing the plants to produce a certain protein in the plant cells. Thus, the present invention has been completed.

That is, the present invention provides:

- A method for giving weed control compoundresistance to a plant which comprises the steps of:
- introducing a gene encoding a protein having the following characteristics (a) to (c):
- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of 25 modifying a substance for which said protein has a specific

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affinity, and

(c) being substantially free from framework regions of variable regions in an immunoglobulin,

into a plant cell; and

expressing the gene (hereinafter referred to as the first aspect of the method of the present invention).

- 2. The method according to the above 1, wherein the gene is introduced into the plant cell in the form that it is operably ligated to a promoter and a terminator both of which are functional in the plant cell.
- 3. The method according to the above 1 or 2, wherein the substance which is concerned with the weed control activity of the weed control compound is the weed control compound itself.
- 4. The method according to the above 1 or 2, wherein the substance which is concerned with the weed control activity of a weed control compound is an endogenous substance in a plant.
- 5. The method according to the above 1 or 2, wherein the weed control compound is that inhibiting porphyrin biosynthesis of a plant.
 - 6. The method according to the above 1 or 2, wherein the weed control compound is a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound.
 - 7. The method according to the above 5 or 6,

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wherein the substance which is concerned with the weed control activity of the weed control compound is protoporphyrin IX.

- 8. The method according to the above 5 or 6, wherein the protein is protoporphyrin IX binding subunit protein of magnesium chelatase, or a variant of said protein having a specific affinity for protoporphyrin IX.
- 9. The method according to the above 8, wherein the protein is magnesium chelatase derived from a photosynthetic microorganism.
- 10. The method according to the above 8, wherein the protein is magnesium chelatase derived from a plant.
- 11. The method according to the above 8, wherein the protein is magnesium chelatase derived from tobacco.
- 12. The method according to the above 5 or 6, wherein the protein comprises the amino acid sequence of SEQ TD NO: 53.
- 13. The method according to the above 5 or 6, wherein the protein has the amino acid sequence of SEQ ID $_{
 m NO}$: 54.
- \$14.\$ The method according to the above 5 or 6, wherein the protein comprises the amino acid sequence of SEQ ID NO: 55.
- 15. The method according to the above 5 or 6, 25 wherein the protein has the amino acid sequence of SEQ ID

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NO: 56.

- \$16.\$ The method according to the above 5 or 6, wherein the protein comprises the amino acid sequence of SEQ ID NO: 57.
- 17. The method according to the above 5 or 6, wherein the protein has the amino acid sequence of SEQ ID NO: 58.
- 18. The method according to the above 5 or 6, wherein the protein comprises the amino acid sequence of SEQ ID NO: 59.
- 19. The method according to the above 5 or 6, wherein the protein has the amino acid sequence of SEQ ID NO: 60.
- \$20.\$ The method according to the above 5 or 6, wherein the protein is composed of 4 to 100 amino acids.
- 21. The method according to the above 5 or 6, wherein the substance which is concerned with the weed control activity of the weed control compound is protoporphyrinogen IX.
- 22. The method according to the above 5 or 6, wherein the protein is a variant of protoporphyrinogen IX oxidase having no capability of oxidizing protoporphyrinogen IX and having a specific affinity for a protoporphyrinogen IX.
- 23. The method according to the above 5 or 6,

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wherein the protein is a variant of protoporphyrinogen IX oxidase having no capability of oxidizing protoporphyrinogen IX and having a specific affinity for a protoporphyrin IX oxidase inhibitory-type herbicidal compound.

- 24. The method according to the above 22 or 23, wherein the protein is a variant of protoporphyrinogen IX oxidase derived from a plant.
- 25. The method according to the above 22 or 23, wherein the protein is a variant of protoporphyrinogen IX oxidase derived from soybean.
- 26. The method according to the above 22 or 23, wherein the protein is a variant of protoporphyrinogen IX oxidase derived from an algae.
- 27. The method according to the above 22 or 23, wherein the protein is a variant of protoporphyrinogen IX oxidase derived from Chlamydomonas.
- 28. A method for giving weed control compoundresistance to a plant which comprises the steps of:

introducing a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for protoporphyrin IX,
- (b) having substantially no capability of modifying protoporphyrinogen IX, and

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(c) being substantially free from framework regions of variable regions in an immunoglobulin, into a plant cell; and

expressing the gene (hereinafter referred to as the second aspect of the method of the present invention).

- 29. The method according to the above 28, wherein the gene is introduced in the plant cell in the form that it is operably ligated to a promoter and a terminator both of which are functional in the plant cell.
- 30. The method according to the above 28 or 29, wherein the weed control compound is that inhibiting porphyrin biosynthesis of a plant.
- 31. The method according to the above 28 or 29, wherein the weed control compound is a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound.
- 32. The method according to the above 30 or 31, wherein the protein is magnesium chelatase or a variant of said protein having a specific affinity for protoporphyrin IX.
- 33. The method according to the above 30 or 31, wherein the protein is ferrochelatase or a variant of said protein having an specific affinity for protoporphyrin IX.
- 34. The method according to the above 30 or 31,wherein the protein is ferrochelatase derived from a plant.

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- 35. The method according to the above 30 or 31, wherein the protein is ferrochelatase derived from barley.
- 36. The method according to the above 30 or 31, wherein the protein is ferrochelatase derived from cucumber.
- $$\,$ 37. The method according to the above 30 or 31, wherein the protein is a peptide composed of 4 to 100 amino acids.
- 38. A method for giving weed control compoundresistance to a plant which comprises the steps of:

introducing a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for protoporphyrinogen IX,
- (b) having the capability for modifying coproporphyrinogen III, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin, into a plant cell; and

 $\mbox{expressing the gene (hereinafter referred to as} \\ \mbox{the third aspect of the method of the present invention).}$

- 39. The method according to the above 38, wherein the gene is introduced into the plant cell in the form that it is operably ligated to a promoter and a terminator both of which are functional in the plant cell.
 - 40. The method according to the above 38 or 39,

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wherein the protein is coproporphyrinogen III oxidase or a variant of said protein having a specific affinity for protoporphyrinogen IX.

- 41. The method according to the above 38 or 39, wherein the protein is coproporphyrinogen III oxidase derived from a microorganism.
- 42 The method according to the above 38 or 39, wherein the protein is coproporphyrinogen III oxidase derived from Escherichia coli.
- 43. A weed control compound-resistant plant whose resistance is given by the method of the above 1, 2, 28 or 29.
- 44. A weed control compound-resistant plant whose resistance is given by the method of the above 38 or 39.
- 45. A method for protecting a plant which comprises applying the weed control compound to a growth area of the plant of the above 43.
- 46. A method for protecting a plant which comprises applying the weed control compound to a growth area of the plant of the above 44.
- 47. A method for selecting a plant which comprises applying a weed control compound to which the plant of the above 43 is resistant to a growth area of the plant of the above 43 and other plants, and selecting

either plant on the basis of difference in growth between the plants.

- 48. A method for selecting a plant which comprises applying a weed control compound to which the plant of the above 44 is resistant to a growth area of the plant of the above 44 and other plants, and selecting either plant on the basis of difference in growth between the plants.
- 49. The method according to the above 47, wherein the plants are plant cells.
- 50. The method according to the above 48, wherein the plants are plant cells.
- 51. The method according to the above 1 or 2, wherein the weed control compound is a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound selected from the compounds of (1) to (3) below, and the substance which is concerned with the weed control activity of the weed control compound is protoporphyrin IX, protoporphyrinogen IX or a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound:
- (1) chlormethoxynil, bifenox, chlornitrofen (CNP), acifluorfen (5-[2-chloro-4-(trifluoromethyl)phenoxy]-2-nitorobenzoic acid) and its ethyl ester, acifluorfen-sodium, oxyfluorfen (2-chloro-1-(3-ethoxy-4-nitrophenoxy)-4-trifluoromethylbenzene), oxadiazon (3-[2,4-dichloro-5-(1-

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methylethoxy)phenyl]-5-(1,1-dimethylethyl)-1,3,4-oxadiazol-2(3H)-one), 2-[4-chloro-2-fluoro-5-(prop-2-ynyloxy)phenyl]-2,3,4,5,6,7-hexahydro-1H-isoindol-1,3-dione, chlorphthalim (N-(4-chlorophenyl)-3,4,5,6-tetrahydrophtalimide), TNPP-ethyl (ethyl 2-[1-(2,3,4-trichlorophenyl)-4-nitropyrazolyl-5-oxy]propionate), or N3-(1-phenylethyl)-2,6-dimethyl-5-propyonylnicotinamide;

(2) a compound represented by the general formula: J-G (I), wherein G is a group represented by any one of the following general formulas G-1 to G-9 and J is a group represented by any one of the following general formulas of J-1 to J-30:

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$$R^{14}$$
 R^{14}
 R^{15}
 R^{14}
 R^{15}
 R

wherein the dotted lines in the formulas J-5, J-6, J-12 and J-24 represent that the left hand ring contains only single bonds, or one bond in the ring is a double bond between carbon atoms;

X is oxygen atom or sulfur atom;

Y is oxygen atom or sulfur atom;

R1 is hydrogen atom or halogen atom;

 $R^2 \quad is \quad hydrogen \quad atom, \quad C_1-C_8alkyl \quad group, \quad C_1-C_8 \\ haloalkyl \quad group, \quad halogen \quad atom, \quad OH \quad group, \quad OR^{27} \quad group, \quad SH \\ group, \quad S(O)_pR^{27} \quad group, \quad COR^{27} \quad group, \quad CO_2R^{27} \quad group, \quad C(O) SR^{27} \\ group, \quad C(O) NR^{29}R^{30} \quad group, \quad CHO \quad group, \quad CR^{27}=NOR^{36} \quad group, \\ CH=CR^{37}CO_2R^{27} \quad group, \quad CH_2CHR^{37}CO_2R^{27} \quad group, \quad CO_2N=CR^{31}R^{32} \quad group, \\ \end{array}$

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nitro group, cyano group, $NHSO_2R^{33}$ group, $NHSO_2NHR^{33}$ group, $NR^{27}R^{39}$ group, NH_2 group or phenyl group optionally substituted with one or more and the same or different C_1 - C, alkyl groups;

p is 0, 1 or 2;

 $$\rm R^3$$ is $\rm C_1-\rm C_2$ alkyl group, $\rm C_1-\rm C_2$ haloalkyl group, OCH3 group, SCH3 group, OCHF2 group, halogen atom, cyano group or nitro group;

 $$\rm R^4$$ is hydrogen atom, $\rm C_1-\rm C_3$ alkyl group, $\rm C_1-\rm C_3$ haloalkyl group or halogen atom;

 R^5 is hydrogen atom, C_1-C_3 alkyl group, halogen atom, C_1-C_3 haloalkyl group, cyclopropyl group, vinyl group, C_2 alkynyl group, cyano group, $C\left(O\right)R^{36}$ group, CO_2R^{39} group, $C\left(O\right)NR^{38}R^{39}$ group, $CR^{34}R^{35}CO$ group, $CR^{34}R^{35}CO_2R^{36}$ group, $CR^{34}R^{35}CO_2R^{36}$ group, $CR^{34}R^{35}CO_3R^{36}$ group, $CR^{34}R^{35}CO_3R^{36}$ group, $CR^{34}R^{35}CO_3R^{36}$ group, $CR^{34}R^{35}CO_3R^{36}$ group, $CR^{34}R^{35}CO_3R^{36}$ group, or, when G is G-2 or G-6, R^4 and R^5 may form C=0 group together with the carbon atom to which they are attached;

 $$R^6$$ is C_1-C_6 alkyl group, C_1-C_6 haloalkyl group, C_2-C_6 alkoxyalkyl group, C_3-C_6 alkenyl group or C_3-C_6 alkynyl group;

 X^1 is single bond, oxygen atom, sulfur atom, NH group, $N(C_1\text{-}C_3 \text{ alkyl})$ group, $N(C_1\text{-}C_3 \text{ haloalkyl})$ group or N(allyl) group;

 R^7 is hydrogen atom, C_1 - C_6 alkyl group, C_1 - C_6

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haloalkyl group, halogen atom, $S(0)_2(C_1-C_6alkyl)$ group or $C(=0)R^{40}$ group;

 $\rm R^8$ is hydrogen atom, $\rm C_1-\rm C_8$ alkyl group, $\rm C_3-\rm C_8$ cycloalkyl group, $\rm C_3-\rm C_8$ alkenyl group, $\rm C_3-\rm C_8$ alkoxyalkyl group, $\rm C_1-\rm C_8$ haloalkyl group, $\rm C_2-\rm C_8$ alkoxyalkyl group, $\rm C_3-\rm C_8$ alkoxyalkyl group, $\rm C_3-\rm C_8$ alkoxyalkyl group, $\rm C_3-\rm C_8$ haloalkynyl group, $\rm C_3-\rm C_8$ haloalkenyl group, $\rm C_1-\rm C_8$ alkylsulfonyl group, $\rm C_1-\rm C_8$ haloalkylsulfonyl group, $\rm C_1-\rm C_8$ alkoxycarbonylalkyl group, $\rm S(O)_2NH(C_1-C_8$ alkyl) group, $\rm C(O)_2NH(C_1-C_8$ alkyl) group, $\rm C(O)_2NH(C_1-C_8$ alkyl) group, to be substituted with $\rm R^{42}$;

 $\,$ n and m are independently 0, 1, 2 or 3 and m + n is 2 or 3;

 $\label{eq:Z} Z \mbox{ is } CR^9R^{10} \mbox{ group, oxygen atom, sulfur atom, } S(0)$ group, $S(0)_2 \mbox{ group or } N(C_1-C_4 \mbox{ alkyl}) \mbox{ group;}$

each R^9 is independently hydrogen atom, C_1-C_3 alkyl group, halogen atom, hydroxyl group, C_1-C_6 alkoxy group, C_1-C_6 haloalkyl group, C_1-C_6 haloalkyl group, C_2-C_6 alkylcarbonyloxy group or C_2-C_6 haloalkylcarbonyloxy group;

each R^{10} is independently hydrogen atom, $C_1\text{--}C_3$ alkyl group, hydroxyl group or halogen atom;

 R^{11} and R^{12} are independently hydrogen atom, halogen atom, C_1-C_6 alkyl group, C_3-C_6 alkenyl group or C_1-C_6 haloalkyl group;

 $R^{13} \ \ is \ \ hydrogen \ \ atom, \ \ C_1-C_6 \ \ alkyl \ \ group, \ \ C_1-C_6$ 25 haloalkyl group, C_3-C_6 alkenyl group, C_3-C_6 haloalkyl group, group,

 C_3-C_6 alkynyl group, C_3-C_6 haloalkynyl group, HC (=0) group, $(C_1-C_4$ alkyl)C (=0) group or NH₂ group;

 $$R^{14}$$ is C_1-C_6 alkyl group, C_1-C_6 alkylthio group, C_1-C_6 haloalkyl group or $N(CH_3)_2$ group;

W is nitrogen atom or CR15;

 R^{15} is hydrogen atom, C_1-C_6 alkyl group, halogen atom, or phenyl group optionally substituted with C_1-C_6 alkyl group, one or two halogen atoms, C_1-C_6 alkoxy group or CF_3 group;

each Q is independently oxygen atom or sulfur
atom;

Q1 is oxygen atom or sulfur atom;

 $Z^1 \text{ is } CR^{16}R^{17} \text{ group, oxygen atom, sulfur atom, } S(0)$ group, $S(0)_2$ group or $N(C_1-C_4alkyl)$ group;

each R^{16} is independently hydrogen atom, halogen atom, hydroxyl group, C_1 - C_6 alkoxy group, C_1 - C_6 haloalkyl group, C_1 - C_6 haloalkoxy group, C_2 - C_6 alkylcarbonyloxy group or C_2 - C_6 haloalkylcarbonyloxy group;

 $\mbox{ each } R^{17} \mbox{ is independently hydrogen atom, hydroxyl} \\ \mbox{ group or halogen atom;}$

 R^{18} is $C_1 \text{--} C_6$ alkyl group, halogen atom or $C_1 \text{--} C_6$ haloalkyl group;

 $$R^{19}$$ and $$R^{20}$$ are independently hydrogen atom, $C_1\text{--}C_6$ alkyl group, or $C_1\text{--}C_6$ haloalkyl group;

 Z^2 is oxygen atom, sulfur atom, NR 9 group or CR 9 R 10

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group;

 $$R^{21}$$ and $$R^{22}$$ are independently C_1-C_6 alkyl group, C_1-C_6 haloalkyl group, C_3-C_6 alkenyl group, C_3-C_6 haloalkenyl group, C_3-C_6 alkynyl group or C_3-C_6 haloalkynyl group;

 \mathbb{R}^{23} is hydrogen atom, halogen atom or cyano group;

 R^{24} is C_1-C_6 alkylsulfonyl group, C_1-C_6 alkyl group, C_3-C_6 haloalkyl group, C_3-C_6 alkenyl group, C_3-C_6 alkynyl group, C_1-C_6 alkoxy group, C_1-C_6 haloalkoxy group or halogen atom;

 $R^{25} \mbox{ is } C_1^-C_6 \mbox{ alkyl group, } C_1^-C_6 \mbox{ haloalkyl group, } C_3^-C_6 \mbox{ alkenyl group or } C_3^-C_6 \mbox{ alkynyl group;}$

 R^{26} is C_1-C_6 alkyl group, C_1-C_6 haloalkyl group or phenyl group optionally substituted with C_1-C_6 alkyl, one or two halogen atoms, one or two nitro groups, C_1-C_6 alkoxy group or CF₃ group;

W1 is nitrogen atom or CH group;

\$T\$ is a group represented by any one of the following general formulas T-1, T-2 and T-3;

(wherein E^1 , E^2 , E^3 , E^4 , E^5 , E^6 , E^7 , E^8 , E^9 , E^{10} , E^{11} and E^{12} are

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independently hydrogen atom or C1-C3 alkyl group);

R27 is C1-Ca alkyl group, C3-Ca cycloalkyl group, C3-C8 alkenyl group, C3-C8alkynyl group, C1-C8 haloalkyl group, C2-C8 alkoxyalkyl group, C2-C8 alkylthioalkyl group, C2-C8 alkylsulfinylalkyl group, C2-C8 alkylsulfonylalkyl group, C1-C8 alkylsulfonyl group, phenylsulfonyl group whose phenyl ring may be substituted with at least one substituent selected from the group consisting of halogen atom and C,alkyl group, C₄-C₈ alkoxyalkoxyalkyl group, C₄-C₈ cycloalkylalkyl group, C_6-C_8 cycloalkoxyalkyl group, C_4-C_8 alkenyloxyalkyl group, C4-C8 alkynyloxyalkyl group, C3-C8 haloalkoxyalkyl group, C4-C8 haloalkenyloxyalkyl group, C4-C8 haloalkynyloxyalkyl group, Cs-Ca cycloalkylthioalkyl group, C.-C. alkenylthioalkyl group, C.-C. alkynylthioalkyl group, C1-C4 alkyl group substituted with phenoxy group whose ring is substituted with at least one substituent selected from the group consisting of halogen atom, C1-C3 alkyl group and C1-C3 haloalkyl group, benzyloxy group whose ring is substituted with at least one substituent selected from the group consisting of halogen atom, C1-C3 alkyl group and C1-C3 haloalkyl group, C4-C8 trialkylsilylalkyl group, C3-C8 C.-C. halocycloalkyl cvanoalkyl group, group, C3-C8 C5-C8 alkoxyalkenyl haloalkenvl group, group, Cs-Cs haloalkoxyalkenyl group, Cs-Cs alkylthioalkenyl group, C3-C8 haloalkynyl group, C5-C8 alkoxyalkynyl group, C5-C8

haloalkoxyalkynyl group, C_5-C_8 alkylthioalkynyl group, C_2-C_8 alkylcarbonyl group, benzyl group whose ring is substituted with at least one substituent selected from the group consisting of halogen atom, C_1-C_3 alkyl group and C_1-C_3 haloalkyl group, $CHR^{34}COR^{28}$ group, $CHR^{34}COOR^{28}$ group,

 $R^{28} \mbox{ is } C_1-C_6 \mbox{ alkyl group, } C_2-C_6 \mbox{ alkenyl group, } C_3-C_6 \mbox{ alkynyl group or tetrahydrofuranyl group;}$

 $$R^{29}$$ and $$R^{31}$$ are independently hydrogen atom or $C_1\text{--}$ C_4 alkyl group;

 R^{30} and R^{32} are independently $C_1\text{-}C_4$ alkyl group or phenyl group whose ring may be substituted with at least one substituent selected from the group consisting of halogen atom, $C_1\text{-}C_3$ alkyl group and $C_1\text{-}C_3$ haloalkyl group; or,

 $$R^{29}$$ and $$R^{10}$$ together may form $-(CH_2)_5-$, $-(CH_2)_4-$ or $-CH_2CH_2CH_2CH_2-$, or the ring thus formed may be substituted with at least one substituent selected from the group consisting of C_1-C_3 alkyl group, phenyl group and benzyl group; or,

 R^{31} and R^{32} may from C_3-C_8 cycloalkyl group together with the carbon atom to which they are attached;

 $R^{33} \mbox{ is } C_1 - C_4 \mbox{ alkyl group, } C_1 - C_4 \mbox{ haloalkyl group;}$ $C_1 - C_6 \mbox{ alkenyl group;}$

 R^{34} and R^{35} are independently hydrogen atom or C_1 -

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C4 alkyl group;

 $R^{36} \ \ is \ \ hydrogen \ \ atom, \ \ C_1-C_6 \ alkyl \ \ group, \ \ C_3-C_6$ alkenyl group or C_3-C_6 alkynyl group;

 R^{37} is hydrogen atom, $C_1 - C_4$ alkyl group or halogen atom;

 R^{36} is hydrogen atom, C_1-C_6 alkyl group, C_3-C_6 cycloalkyl group, C_3-C_6 alkenyl group, C_3-C_6 alkoxyalkyl group, C_1-C_6 haloalkyl group, phenyl group whose ring may be substituted with at least one substituent selected from the group consisting of halogen atom, C_1-C_4 alkyl group and C_1-C_4 alkoxy group, $-CH_2CO_2(C_1-C_4$ alkyl) group or $-CH_1CH_3)CO_2(C_1-C_4$ alkyl) group;

 $R^{39} \quad \text{is hydrogen atom,} \quad C_1-C_2 \quad \text{alkyl group of } C(0)O(C_1-C_4 \text{ alkyl) group;}$

 $R^{40} \ \ is \ \ hydrogen \ \ atom, \ \ C_1-C_6 \ \ alkyl \ \ group, \ \ C_1-C_6$ alkovy group or NH (C_1-C_6 alkyl) group;

 R^{41} is $C_1{}^-C_6$ alkyl group, $C_1{}^-C_6$ haloalkyl group, $C_1{}^ C_6$ alkoxy group, NH(C_1{}^-C_6 alkyl) group, phenyl group whose ring may be substituted with one substituent selected from the group consisting of R^{42} group, benzyl group and $C_2{}^-C_8$ dialkylamino group; and

 $R^{42} \text{ is } C_1\text{--}C_6 \text{ alkyl group, one or two halogen atoms,} \\ C_1\text{--}C_6 \text{ alkoxy group or } CF_3 \text{ group;}$

(3) a compound of the formula (II):

or nipilacrofen,

wherein R43 is C1-C4 alkyl group;

 R^{44} is C_1 - C_4 alkyl group, C_1 - C_4 alkylthio group, C_1 - C_4 alkoxy group, C_1 - C_4 haloalkyl group, C_1 - C_4 haloalkylthio group or C_1 - C_4 haloalkoxy group;

 R^{43} and R^{44} together may form $-(CH_2)_3-$ or $-(CH_2)_4-$; R^{45} is hydrogen atom or halogen atom; $R^{46} \text{ is hydrogen atom or } C_1-C_4 \text{ alkyl group;}$ $R^{47} \text{ is hydrogen atom, nitro group, cyano group,}$

R⁴⁷ is hydrogen atom, nitro group, cyano gro -COOR⁴⁹ group, -C(=X)NR³⁰R⁵¹ group or -C(=X²)R⁵² group;

R⁴⁸ is hydrogen atom, halogen atom, cyano group,
C₁-C₄ alkyl group optionally substituted with at least one
substituent selected from the group consisting of halogen
15 atom and hydroxyl group, C₁-C₄ alkoxy group, phenyl group
optionally substituted with at least one substituent
selected from the group consisting of halogen atom, nitro
group, cyano group, C₁-C₄ alkyl group, C₁-C₄ alkoxy group and
halo-C₁-C₄ alkyl group, pyrrolyl group, C₂-C₆ alkyl group,
C₃-C₆ alkenyl group, C₃-C₆ alkynyl group, C₅-C₆ alkoxy group,
a group selected from the group consisting of C₂-C₆ alkyl

group, C_3-C_8 alkenyl group, C_3-C_8 alkynyl group and C_3-C_8 alkoxy group into which at least one oxygen atom is inserted, or any one of groups represented by the following formulas:

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wherein $R^{19},\ R^{50}$ and R^{52} are, the same or different, hydrogen atom or C_1-C_4 alkyl group;

 $$R^{50}$$ and $$R^{51}$$ may form saturated alicyclic 5 or 6 membered ring together with the nitrogen atom to which they are attached;

 $$R^{52}$$ is hydrogen atom, $C_1\text{-}C_4$ alkyl group or $C_1\text{-}C_4$ alkyl group substituted with at least one halogen atom;

 R^{33} is hydrogen atom, C_1 - C_4 alkyl group optionally substituted with at least one halogen atom, C_2 - C_6 alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl group optionally substituted with at least one halogen atom, phenyl group optionally substituted with at least one halogen atom, C_3 - C_8 cycloalkyl group, cyanomethyl group, or R^{63} CO- group;

 R^{54} is hydrogen atom, C_1 - C_6 alkyl group optionally substituted with at least one halogen atom, C_2 - C_6 alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl group optionally substituted with at least one halogen atom, phenyl group optionally substituted with halogen atom, C_3 - C_6 cycloalkyl group, cyanomethyl group, C_1 - C_4 alkoxy- C_1 - C_6 alkyl group, di- C_1 - C_4 alkylamino- C_1 - C_4 alkyl group, tetrahydrofurfurylmethyl group, C_3 - C_6 alkynyloxy- C_1 - C_6 alkyl group, benzyl whose ring may be substituted with substituent selected from the group consisting of halogen atom, nitro group, cyano group, C_1 - C_4 alkyl group, C_1 - C_6

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alkoxy group and halo- C_1 - C_4 alkyl group, $-C(=X^2)\,R^{63}$ group, - $(CH_2)_a - (O)_d - R^{70}$ group, $-(CH_2)_a - O - (CH_2)_b - R^{70}$ group, $-(CH_2)_a - X^2 - R^{76}$ group;

R53 and R54 together with the nitrogen atom to which they are attached may form saturated alicyclic 3, 5 or 6 membered ring or aromatic 5 or 6 membered ring in which a carbon atom may be optionally replaced with oxygen atom;

R55 is hydrogen atom, C1-C4 alkyl group, C2-C6 alkenyl group or C3-C6 alkynyl group, or R55 and R56 together may form - (CH2)e-;

R⁵⁶ and R⁵⁷ are independently C₁-C₄ alkyl group optionally substituted with at least one halogen atom, C2-C. alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl optionally substituted with at least one halogen atom or phenyl group optionally substituted with at least one halogen atom, hydrogen atom, C3-C6 cycloalkyl group, -XR60 group or -NR61R62 group;

R58 is hydrogen atom, C1-C6 alkyl group, C2-C6 alkenyl group, C3-C6 alkynyl group, C1-C4 alkylcarbonyl group, cyano-C1-C3 alkyl group, C1-C4 alkoxycarbonyl-C1-C4 alkyl group, di-C1-C4 alkoxycarbonyl-C1-C4 alkyl group, benzyl group, C1-C2 alkoxy-C1-C4 alkynyl group, -(CH2)a-R75 group, - $(CH_2)_a - X^2 - R^{72}$ group, $-(CH_2)_a - X^2 - (CH_2)_b - R^{72}$ group or $-(CH_2)_a - X^2 - (CH_2)_a - R^{72}$ (CH₂)_p-X²-(CH₂)_c-R⁷² group;

 $$R^{59}$$ is hydrogen atom, $C_1\text{-}C_4$ alkyl group, $C_2\text{-}C_6$ alkenyl group, $C_3\text{-}C_6$ alkynyl group, cyano- $C_1\text{-}C_3$ alkyl group, $C_1\text{-}C_4$ alkylcarbonyl- $C_1\text{-}C_3$ alkyl group or phenyl group;

 R^{60} is C_1 - C_4 alkyl group optionally substituted with at least one halogen atom;

 $R^{\varepsilon t}$ and $R^{\varepsilon 2}$ are, the same or different, hydrogen atom or C,-C, alkyl group;

 R^{63} is C_1-C_4 alkyl group optionally substituted with at least one halogen atom, C_1-C_4 alkoxy- C_1-C_4 alkyl group, C_3-C_6 cycloalkyl group, C_1-C_4 alkylthio- C_1-C_4 alkyl group, C_3-C_6 cycloalkyl group, phenyl group whose ring may be substituted with one substituent selected from the group consisting of halogen atom, nitro group, cyano group, C_1-C_4 alkyl group, C_1-C_4 alkoxy group and halo- C_1-C_4 alkyl group, $-NR^{73}R^{74}$ group or $-(CH_2)_3-(O)_4-R^{75}$ group;

 $\label{eq:resolvent} R^{64} \quad \text{is} \quad C_1 - C_4 \quad \text{alkoxycarbonyl} \quad \text{group} \quad \text{or} \quad \text{carboxyl} \\ \text{group;}$

 R^{65} is chloromethyl group, cyanomethyl group, C_3 - C_6 cycloalkyl group into which at least one oxygen atom may be inserted, or C_3 - C_4 alkoxycarbonyl- C_3 - C_4 alkyl group;

 R^{66} is hydroxyl group or $-NR^{67}R^{68}$ group;

A is $-NR^{67}R^{68}$ group or $-S(0)_f-R^{69}$ group;

 $$R^{67}$$ and $$R^{68}$$ are, the same or different, hydrogen atom or C,-C, alkyl group;

R⁶⁹ is C₁-C₄ alkyl group or C₁-C₄ haloalkyl group;

 R^{70} is hydrogen atom, hydroxyl group, halogen atom, C_1 - C_4 alkyl group optionally substituted with at least one C_1 - C_4 alkoxy group, C_3 - C_6 cycloalkyl group into which at least one oxygen atom may be inserted, C_3 - C_6 cycloalkyl group optionally substituted with one or two methyl groups, furyl group, thienyl group or -C (=0) R^{71} group;

 R^{71} and R^{72} are, the same or different, $C_1\!-\!C_4$ alkyl group or $C_1\!-\!C_4$ alkoxy group;

 $$R^{73}$$ and $$R^{74}$$ are, the same or different, $C_1\!-\!C_4$ alkyl group or phenyl group;

 R^{75} is C_3-C_6 cycloalkyl into which at least one oxygen atom may be inserted, C_3-C_6 cycloalkyl group optionally substituted with one or two methyl groups, furyl group, thienyl group or -C (=0) R^{71} group;

R76 is C1-C4 alkyl group;

a, b and c is independently 1, 2 or 3;

d is 0 or 1:

e is 2 or 3;

f is 1 or 2; and

X2 is oxygen atom or sulfur atom,

52. The method according to the above 1, additionally comprising the steps of:

introducing into the plant cell, a second gene selected from a gene encoding a protein substantially having protoporphyrinogen oxidase activity, a gene encoding

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a protein substantially having 5-enolpyruvylshikamate-3phosphate synthase activity and a gene encoding a protein substantially having glyphosate oxidoreductase activity; and

expressing said second gene.

53. A plant cell having:

a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin; and

at least one altered form of an enzymatic activity which gives a resistance to a weed control compound in an amount inhibiting a naturally occurring form of said enzymatic activity, wherein said altered form of an enzymatic activity is a form of enzymatic activity selected from a protoporphyrinogen oxidase activity, 5-enolpyruvylshikamate-3-phosphate synthase activity and glyphosate oxidoreductase activity.

54. A plant cell having:

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a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin; and an altered protoporphyrinogen oxidase activity which gives a resistance to a weed control compound in an amount inhibiting a natural occurring protoporphyrinogen
 - 55. A plant cell having:

oxidase activity.

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from frameworkregions of variable regions in an immunoglobulin; and

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an altered 5-enolpyruvylshikamate-3-phosphate synthase activity which gives a resistance to a weed control compound in an amount inhibiting a natural occurring 5-enolpyruvylshikamate-3-phosphate synthase activity.

56. A plant cell having:

a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- $\mbox{(b) having substantially no capability of} \\ \mbox{modifying a substance for which said protein has a specific} \\ \mbox{affinity, and} \\$

(c) being substantially free from framework

- regions of variable regions in an immunoglobulin; and

 an altered glyphosate oxidoreductase activity
 which gives a resistance to a weed control compound in an
 amount inhibiting a natural occurring glyphosate
 oxidoreductase activity.
- 57. The plant cell according to the above 53, wherein said altered form of an enzymatic activity is conferred by a second gene selected from a gene encoding a protein substantially having a protein substantially having 5-

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enolpyruvylshikamate-3-phosphate synthase activity and a gene encoding a protein substantially having glyphosate oxidoreductase activity.

- 58. The plant cell according to the above 57, wherein the gene encoding a protein having the following characteristics (a) to (c):
- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin; and

the second gene are introduced into the plant cell in the form in that both of said genes are operably ligated to a promoter and a terminator both of which are functional in said plant cell.

59. The plant cell according to the above 57, wherein the protein substantially having a protoporphyrinogen IX oxidase activity is protoporphyrinogen IX oxidase, the protein substantially having a 5-enolpyruvylshikamate-3-phosphate synthase activity is 5-enolpyruvylshikamate-3-phosphate synthase and the protein substantially having glyphosate oxidoreductase activity is

glyphosate oxidoreductase.

- 60. The plant cell according to the above 53, wherein the plant cell is derived from dicotyledones or monocotyledones.
- 61. A plant comprising the plant cell of the above 54.
 - $\ensuremath{\text{62.}}$ A plant comprising the plant cell of the above 55.
 - 63. A plant comprising the plant cell of the above 56.
 - 64. A method for protecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound to a growth area of the plant of the above 61.
 - 65. A method for protecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of the above 62.
- 20 66. A method for protecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of the above 63.
 - 67. A method for selecting a plant which

comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound to a growth area of the plant of the above 61 and other plants, and selecting either plant on the basis of difference in growth between the plants.

- 68. A method for selecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of the above 62 and other plants, and selecting either plant on the basis of difference in growth between the plants.
- 69. A method for selecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of the above 63 and other plants, and selecting either plant on the basis of difference in growth between the plants.

DETAILED DESCRIPTION OF THE INVENTION

In the method of the present invention, substances which are concerned with weed control activities of weed control compounds (hereinafter referred to as weed control substances) are those constituting a part of metabolic reaction systems in organisms which are

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responsible for weed control activities upon applying the compounds to plants. Examples thereof include weed control compounds themselves, endogenous substances in plants, and the like. Specifically, as such endogenous substances in plants, for example, there are substrates of target enzymes on which weed control compounds act, or precursors or metabolites of the substrates which cause cellular dysfunction upon accumulating in plant cells; substances produced by the above substances in plant cells which cause cellular dysfunction; and the like. More specifically, it has been known that, when a compound having herbicidal activity (hereinafter referred to as herbicidal compound) which inhibits the activity of protoporphyrinogen oxidase (EC 1.3.3.4, hereinafter referred to as PPO) applied to a plant, protoporphyrinogen IX which is the substrate of PPO is accumulated in the plant cells and it is metabolized to form protoporphyrin X, followed by formation of active oxygen in the presence of both protoporphyrin X and light in the cells, which damages cell functions (Junshi MIYAMOTO ed., Atarashii Noyaku no Kagaku (Chemistry of New Agrochemicals), Chapter 3, Section 3.3, p Hirokawa 106 (1993). Shoten, Tokyo]. protoporphyrinogen IX, protoporphyrin IX and active oxygen in these systems, and the like can be exemplified as these substances.

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In the method of the present invention, weed control compounds include compounds having herbicidal activities, plant growth regulator activities, and the like.

Examples of the herbicidal compounds include compounds inhibiting porphyrin biosynthesis, compounds inhibiting electron transfer in photosynthesis, compounds inhibiting carotenoid biosynthesis, compounds inhibiting acid biosynthesis, compounds inhibiting biosynthesis, compounds inhibiting cell wall biosynthesis, compounds influencing protein biosynthesis, nucleic acid biosynthesis and cell division, compounds having auxin antagonistic activity, and the like. More specifically, as inhibiting porphyrin biosynthesis, for compounds example, there are compounds inhibiting PPO activity (PPO inhibitory-type herbicidal compound), and the like. As the compounds inhibiting electron transfer in photosynthesis, for example, there are compounds inhibiting electron transfer of photochemical system I or II, compounds 4-hydroxyphenyl pyruvate dioxygenase (EC inhibiting 1.13.11.27; hereinafter referred to as 4-HPPD) which influences biosynthesis of plastoquinone which transfers electrons, and the like. As the compounds inhibiting carotenoid biosynthesis, for example, there are compounds inhibiting phytoene desaturase (hereinafter referred to as PDS), and the like. As the compounds inhibiting amino acid

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biosynthesis, for example, there are compounds inhibiting EPSPS, acetolactate synthase (EC 4.1.3.18; hereinafter referred to as ALS), glutamine synthetase (EC 6.3.1.2; hereinafter referred to as GS), dihydropteroate synthase (EC 2.5.1.15; hereinafter referred to as DHP), and the like. As the compounds inhibiting lipid biosynthesis, for example, there are compounds inhibiting acetyl CoA carboxylase (EC 6.4.1.2; hereinafter referred to as ACC), and the like. As compounds inhibiting cell wall biosynthesis, the compounds inhibiting cellulose example, there are biosynthesis, and the like. As the compounds influencing protein biosynthesis, nucleic acid biosynthesis or cell division, for example, there are compounds inhibiting formation of microtubules, and the like.

Examples of the compounds having plant growth regulator activities include compounds having antagonistic activities against plant hormones which enhance cell elongation and differentiation, and the like. Specifically, for example, there are 2,4-D, phenoxyalkane carboxylic acid, derivatives of benzoic acid, derivatives of picolinic acid, and the like.

As the above-described PPO inhibitory-type herbicidal compounds, for example, there are the compounds disclosed in Duke, S.O., Rebeiz, C.A., ACS Symposium Series 559, Porphyric Pesticides, Chemistry, Toxicology, and

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Pharmaceutical Applications, American Chemical Society, Washington DC (1994), and the like. Specifically, examples thereof include the following compounds:

- (1) chlormethoxynil, bifenox, chlornitrofen (CNP), (5-[2-chloro-4-(trifluoromethyl)phenoxy]-2acifluorfen nitorobenzoic acid) and its ethyl ester, acifluorfen-sodium, (2-chloro-1-(3-ethoxy-4-nitrophenoxy)-4oxyfluorfen (3-[2,4-dichloro-5-(1oxadiazon trifluorobenzene), methylethoxy)phenyl]-5-(1,1-dimethylethyl)-1,3,4-oxydiazol-2-[4-chloro-2-fluoro-5-(prop-2-2-(3H)-one), ynyloxy)phenyl]-2,3,4,5,6,7-hexahydro-1H-isoindol-1,3-dione, (N-(4-chlorophenyl)-3,4,5,6chlorphthalim, TNPP-ethyl (ethyl 2-[1-(2,3,4tetrahydrophtalimide), trichlorophenyl)-4-nitropyrazolyl-5-oxy]propionate), or N3-(1-phenylethyl)-2,6-dimethyl-5-propyonylnicotinamide;
- (2) a compound represented by the general fomrula: J-G (I), wherein G is a group represented by any one of the following general formulas G-1 to G-9 and J is a group represented by any one of the following general formulas J-1 to J-30:

J-12

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wherein the dotted lines in the formulas J-5, J-6, J-12 and J-24 represent that the left hand ring contains only single bonds, or one bond in the ring is a double bond between carbon atoms;

X is oxygen atom or sulfur atom;

Y is oxygen atom or sulfur atom;

R1 is hydrogen atom or halogen atom;

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substituted with one or more and the same or different C_1 - C_4 alkyl groups;

p is 0, 1 or 2;

R³ is C₁-C₂ alkyl group, C₁-C₂ haloalkyl group, OCH₃ group, SCH₃ group, OCHF₂ group, halogen atom, cyano group or nitro group;

 $R^4 \quad \mbox{is hydrogen atom,} \quad C_1-C_3 \quad \mbox{alkyl group,} \quad C_1-C_3$ haloalkyl group or halogen atom;

 R^5 is hydrogen atom, C_1-C_3 alkyl group, halogen atom, C_1-C_3 haloalkyl group, cyclopropyl group, vinyl group, C_2 alkynyl group, cyano group, $C\left(O\right)R^{38}$ group, CO_2R^{18} group, $C\left(O\right)NR^{38}R^{39}$ group, $CR^{34}R^{35}CO$ group, $CR^{24}R^{35}CO_2R^{38}$ group, $CR^{34}R^{35}CO_2R^{38}$ group, $CR^{34}R^{35}CO_2R^{38}$ group, $CR^{34}R^{35}CO_2R^{38}$ group, $CR^{34}R^{35}CO_2R^{38}$ group, or OCHR $^{34}OC\left(O\right)NR^{38}R^{39}$ group, or, when G is G-2 or G-6, R^4 and R^5 may form C=O group together with the carbon atom to which they are attached;

 R^6 is C_1-C_6 alkyl group, C_1-C_6 haloalkyl group, C_2-C_6 alkoxyalkyl group, C_3-C_6 alkenyl group or C_3-C_6 alkynyl group;

 X^1 is single bond, oxygen atom, sulfur atom, NH group, $N(C_1-C_3$ alkyl) group, $N(C_1-C_3$ haloalkyl) group or N(allyl) group;

 R^7 is hydrogen atom, C_1-C_ϵ alkyl group, C_1-C_ϵ haloalkyl group, halogen atom, $S\left(O\right)_2\left(C_1-C_\epsilon alkyl\right)$ group or $C\left(=O\right)R^{40}$ group;

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 $\rm R^8$ is hydrogen atom, $\rm C_1-\rm C_8$ alkyl group, $\rm C_3-\rm C_8$ cycloalkyl group, $\rm C_3-\rm C_8$ alkenyl group, $\rm C_3-\rm C_8$ alkynyl group, $\rm C_1-\rm C_8$ haloalkyl group, $\rm C_2-\rm C_8$ alkoxyalkyl group, $\rm C_3-\rm C_8$ alkoxyalkyl group, $\rm C_3-\rm C_8$ alkoxyalkyl group, $\rm C_3-\rm C_8$ haloalkynyl group, $\rm C_3-\rm C_8$ haloalkynyl group, $\rm C_1-\rm C_8$ haloalkylsulfonyl group, $\rm C_1-\rm C_8$ haloalkylsulfonyl group, $\rm C_1-\rm C_8$ alkoxycarbonylalkyl group, $\rm S(O)_2NH(C_1-C_8$ alkyl) group, $\rm C(O)\,R^{41}$ group or benzyl group whose phenyl ring may be substituted with $\rm R^{42}$;

n and m are independently 0, 1, 2 or 3 and m + n is 2 or 3:

 $Z \ \ is \ CR^9R^{10} \ group, \ oxygen \ atom, \ sulfur \ atom, \ S(O)$ group, $S(O)_2$ group or $N(C_1-C_4 \ alkyl)$ group;

each R^9 is independently hydrogen atom, C_1-C_3 alkyl group, halogen atom, hydroxyl group, C_1-C_6 alkoxy group, C_1-C_6 haloalkyl group, C_1-C_6 haloalkyl group, C_2-C_6 alkylcarbonyloxy group or C_2-C_6 haloalkylcarbonyloxy group;

each R^{10} is independently hydrogen atom, C_1-C_3 alkyl group, hydroxyl group or halogen atom;

 R^{11} and R^{12} are independently hydrogen atom, halogen atom, C_1-C_6 alkyl group, C_3-C_6 alkenyl group or C_1-C_6 haloalkyl group;

 R^{13} is hydrogen atom, C_1-C_6 alkyl group, C_1-C_6 haloalkyl group, C_3-C_6 alkenyl group, C_3-C_6 haloalkenyl group, C_3-C_6 alkynyl group, C_3-C_6 haloalkynyl group, C_3-C_6 haloalkynyl group, C_3-C_6 haloalkynyl group, C_3-C_6 alkyl) C_3-C_6 group or NH_2 group;

 $R^{14} \text{ is } C_1^-C_6 \text{ alkyl group, } C_1^-C_6 \text{ alkylthio group, } C_1^-C_6 \text{ haloalkyl group or } N(CH_3)_2 \text{ group;}$

W is nitrogen atom or CR15;

 R^{15} is hydrogen atom, C_1-C_6 alkyl group, halogen atom, or phenyl group optionally substituted with C_1-C_6 alkyl group, one or two halogen atoms, C_1-C_6 alkoxy group or CF, group;

 $\mbox{ each Q is independently oxygen atom or sulfur} \\ \mbox{atom;}$

 Q^1 is oxygen atom or sulfur atom;

 $Z^1 \text{ is } CR^{16}R^{17} \text{ group, oxygen atom, sulfur atom, } S(0) \\$ group, $S(0)_2 \text{ group or } N(C_1-C_4alkyl) \text{ group;}$

each R^{16} is independently hydrogen atom, halogen atom, hydroxyl group, C_1 - C_6 alkoxy group, C_1 - C_6 haloalkyl group, C_1 - C_6 haloalkoxy group, C_2 - C_6 alkylcarbonyloxy group or C_2 - C_6 haloalkylcarbonyloxy group;

each R^{17} is independently hydrogen atom, hydroxyl group or halogen atom;

 $$R^{18}$$ is $C_1\text{--}C_6$$ alkyl group, halogen atom or $C_1\text{--}C_6$$ haloalkyl group;

 $$R^{10}$$ and $$R^{20}$$ are independently hydrogen atom, $C_1 - C_6$ alkyl group, or $C_1 - C_6$ haloalkyl group;

 Z^2 is oxygen atom, sulfur atom, NR^{ϑ} group or $CR^{\vartheta}R^{10}$ group;

 R^{21} and R^{22} are independently C_1-C_6 alkyl group, C_1-

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 C_6 haloalkyl group, C_3 - C_6 alkenyl group, C_3 - C_6 haloalkenyl group, C_3 - C_6 alkynyl group or C_3 - C_6 haloalkynyl group;

 $\ensuremath{R^{23}}$ is hydrogen atom, halogen atom or cyano group;

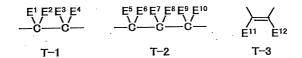
 R^{24} is C_1-C_6 alkylsulfonyl group, C_1-C_6 alkyl group, $C_1-C_6 \text{ haloalkyl group, } C_3-C_6 \text{ alkenyl group, } C_3-C_6 \text{ alkynyl group, } C_1-C_6 \text{ alkoxy group, } C_1-C_6 \text{ haloalkoxy group or halogen atom;}$

 $R^{25} \mbox{ is } C_1-C_6 \mbox{ alkyl group, } C_1-C_6 \mbox{ haloalkyl group, } C_3-C_6 \mbox{ alkenyl group or } C_3-C_6 \mbox{ alkynyl group;}$

 $$R^{26}$$ is C_1-C_6 alkyl group, C_1-C_6 haloalkyl group or phenyl group optionally substituted with C_1-C_6 alkyl, one or two halogen atoms, one or two nitro groups, C_1-C_6 alkoxy group or CF3 group;

W1 is nitrogen atom or CH group;

\$T\$ is a group represented by any one of the following general formulas T-1, T-2 and T-3;



(wherein E^1 , E^2 , E^3 , E^4 , E^5 , E^6 , E^7 , E^8 , E^9 , E^{10} , E^{11} and E^{12} are independently hydrogen atom or C_1-C_3 alkyl group);

 R^{27} is C_1-C_8 alkyl group, C_3-C_8 cycloalkyl group;

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 C_3-C_8 alkenyl group, C_3-C_8 alkynyl group, C_1-C_8 haloalkyl group, C_2-C_8 alkoxyalkyl group, C_2-C_8 alkylthioalkyl group, C_2-C_8 alkylsulfinylalkyl group, C_2-C_8 alkylsulfonylalkyl group, $C_1 - C_8$ alkylsulfonyl group, phenylsulfonyl group whose phenyl ring may be substituted with at least one substituent selected from the group consisting of halogen atom and C_1 alkyl group, C_4-C_8 alkoxyalkoxyalkyl group, cycloalkylalkyl group, C_6-C_8 cycloalkoxyalkyl group, C_4-C_8 alkenyloxyalkyl group, C_4-C_8 alkynyloxyalkyl group, C_3-C_8 haloalkoxyalkyl group, C_4-C_8 haloalkenyloxyalkyl group, C_4-C_8 haloalkynyloxyalkyl group, C_6-C_0 cycloalkylthioalkyl group, C_4-C_8 alkenylthioalkyl group, C_4-C_8 alkynylthioalkyl group, C_1 - C_4 alkyl group substituted with phenoxy group whose ring is substituted with at least one substituent selected from the group consisting of halogen atom, C_1 - C_3 alkyl group and C1-C3 haloalkyl group, benzyloxy group whose ring substituted with at least one substituent selected from the group consisting of halogen atom, C_1-C_3 alkyl group and C_1-C_3 haloalkyl group, C_4 - C_8 trialkylsilylalkyl group, C3-C8 halocycloalkyl group, cyanoalkyl group, C₅-C₈ alkoxyalkenyl group, haloalkenvl group, haloalkoxyalkenyl group, C_5-C_8 alkylthioalkenyl group, C_3-C_8 haloalkynyl group, C₅-C₈ alkoxyalkynyl group, C₅-C₈ haloalkoxyalkynyl group, C_5-C_8 alkylthioalkynyl group, C_2-C_8 alkylcarbonyl group, benzyl group whose ring is substituted

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with at least one substituent selected from the group consisting of halogen atom, C_1 – C_3 alkyl group and C_1 – C_3 haloalkyl group, $CHR^{34}COR^{28}$ group, $CHR^{34}COR^{28}$ group, $CHR^{34}COR^{28}$ group, $CHR^{34}P(S)(OR^{28})_2$ group, $CHR^{34}C(O)NR^{29}R^{30}$ group or $CHR^{34}C(O)NH_3$ group;

 R^{28} is $C_1^-C_6$ alkyl group, $C_2^-C_6$ alkenyl group, $C_3^-C_6$ alkynyl group or tetrahydrofuranyl group;

 $$R^{29}$$ and $$R^{31}$$ are independently hydrogen atom or $C_1\text{--}$ C_4 alkyl group;

 R^{30} and R^{32} are independently $C_1\text{--}C_4$ alkyl group or phenyl group whose ring may be substituted with at least one substituent selected from the group consisting of halogen atom, $C_1\text{--}C_3$ alkyl group and $C_1\text{--}C_3$ haloalkyl group; or,

 R^{29} and R^{30} together may form $-(CH_2)_5-$, $-(CH_2)_4-$ or $-CH_2CH_2OCH_2CH_2-$, or the ring thus formed may be substituted with at least one substituent selected from the group consisting of C_1-C_3 alkyl group, phenyl group and benzyl group; or,

 $$R^{31}$$ and $$R^{32}$$ may from $$C_3\text{-}C_8$$ cycloalkyl group together with the carbon atom to which they are attached;

 $R^{33} \ \mbox{is} \ C_1 - C_4 \ \mbox{alkyl group,} \ C_1 - C_4 \ \mbox{haloalkyl group or}$ $C_3 - C_6 \ \mbox{alkenyl group;}$

 $$\rm R^{34}$$ and $\rm R^{35}$ are independently hydrogen atom or $\rm C_1\textsc{-}C_4$ alkyl group;

 R^{36} is hydrogen atom, C_1-C_6 alkyl group, C_3-C_6

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alkenyl group or C3-C6 alkynyl group;

 $$R^{37}$$ is hydrogen atom, $C_1\!-\!C_4$ alkyl group or halogen atom;

 R^{38} is hydrogen atom, C_1-C_6 alkyl group, C_3-C_6 cycloalkyl group, C_3-C_6 alkenyl group, C_3-C_6 alkoxyalkyl group, C_1-C_6 haloalkyl group, phenyl group whose ring may be substituted with at least one substituent selected from the group consisting of halogen atom, C_1-C_4 alkyl group and C_1-C_4 alkoxy group, $-CH_2CO_2(C_1-C_4$ alkyl) group or $-CH(CH_3)CO_2(C_1-C_4$ alkyl) group;

 $R^{39} \ \mbox{is hydrogen atom,} \ \ C_1-C_2 \ \mbox{alkyl group or}$ $C(0)O(C_1-C_4 \ \mbox{alkyl) group;}$

 $R^{40} \mbox{ is hydrogen atom, } C_1-C_6 \mbox{ alkyl group, } C_1-C_6$ alkoxy group or NH(C1-C6 alkyl) group;

 R^{41} is C_1-C_6 alkyl group, C_1-C_6 haloalkyl group, C_1-C_6 alkoxy group, $NH\left(C_1-C_6$ alkyl) group, phenyl group whose ring may be substituted with one substituent selected from the group consisting of R^{42} group, benzyl group and C_2-C_8 dialkylamino group; and

 $R^{42} \mbox{ is } C_1-C_6 \mbox{ alkyl group, one or two halogen atoms,}$ $C_1-C_6 \mbox{ alkoxy group or } CF_3 \mbox{ group;}$

(3) a compound of the formula (II):

or nipilacrofen,

wherein R43 is C1-C4 alkyl group;

 $R^{44} \ \, \text{is} \ \, C_1-C_4 \ \, \text{alkyl group,} \ \, C_1-C_4 \ \, \text{alkoxy group,} \ \, C_1-C_4 \ \, \text{haloalkyl group,} \ \, C_1-C_4 \ \, \text{haloalkylthio}$ group or $C_1-C_4 \ \, \text{haloalkoxy group;}$

 R^{43} and R^{44} together may form -(CH2) $_3-$ or -(CH2) $_4-$; R^{45} is hydrogen atom or halogen atom;

 R^{46} is hydrogen atom or C_1 - C_4 alkyl group;

 R^{47} is hydrogen atom, nitro group, cyano group, -COOR⁴⁹ group, -C(=X)NR⁵⁰R⁵¹ group or -C(=X²)R⁵² group;

R⁴⁸ is hydrogen atom, halogen atom, cyano group, C₁-C₄ alkyl group optionally substituted with at least one substituent selected from the group consisting of halogen atom and hydroxyl group, C₁-C₄ alkoxy group, phenyl group optionally substituted with at least one substituent selected from the group consisting of halogen atom, nitro group, cyano group, C₁-C₄ alkyl group, C₁-C₄ alkoxy group and halo-C₁-C₄ alkyl group, pyrrolyl group, C₂-C₆ alkyl group, C₃-C₆ alkenyl group, C₃-C₆ alkynyl group, C₃-C₆ alkoxy group, a group selected from the group consisting of C₂-C₆ alkyl group, C₃-C₆ alkenyl group, C₃-C₆ alkenyl group, C₃-C₆ alkynyl group and C₃-C₆ alkoxy group into which at least one oxygen atom is inserted, or any one of groups represented by the following formulas:

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wherein $R^{49},\ R^{50}$ and R^{52} are, the same or different, hydrogen atom or $C_1{-}C_4$ alkyl group;

 $$R^{50}$$ and $$R^{51}$$ may form saturated alicyclic 5 or 6 membered ring together with the nitrogen atom to which they are attached;

 $$R^{52}$$ is hydrogen atom, $C_1\!-\!C_4$ alkyl group or $C_1\!-\!C_4$ alkyl group substituted with at least one halogen atom;

 R^{53} is hydrogen atom, $C_1 - C_4$ alkyl group optionally

substituted with at least one halogen atom, C_2 - C_6 alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl group optionally substituted with at least one halogen atom, phenyl group optionally substituted with at least one halogen atom, C_3 - C_6 cycloalkyl group, cyanomethyl group, or R^6 3CO- group;

 R^{54} is hydrogen atom, C_1-C_6 alkyl group optionally substituted with at least one halogen atom, C_2-C_6 alkenyl group optionally substituted with at least one halogen atom, C_3-C_6 alkynyl group optionally substituted with at least one halogen atom, phenyl group optionally substituted with halogen atom, C_3-C_6 cycloalkyl group, cyanomethyl group, C_1-C_4 alkoxy- C_1-C_6 alkyl group, $di-C_1-C_4$ alkylamino- C_1-C_4 alkyl group, tetrahydrofurfurylmethyl group, C_3-C_6 alkynyloxy- C_1-C_4 alkyl group, benzyl whose ring may be substituted with substituent selected from the group consisting of halogen atom, nitro group, cyano group, C_1-C_4 alkyl group, C_1-C_4 alkoxy group and halo- C_1-C_4 alkyl group, $-C_1-C_4$ alkoxy group and halo- C_1-C_4 alkyl group, $-C_1-C_4$ alkoxy group and halo- C_1-C_4 alkyl group, $-C_1-C_4$ alkyl gro

 ${
m R}^{53}$ and ${
m R}^{54}$ together with the nitrogen atom to which they are attached may form saturated alicyclic 3, 5 or 6 membered ring or aromatic 5 or 6 membered ring in which a carbon atom may be optionally replaced with oxygen

25 atom;

 $$R^{55}$$ is hydrogen atom, C_1-C_4 alkyl group, C_2-C_6 alkenyl group or C_3-C_6 alkynyl group, or R^{55} and R^{56} together may form -(CH,),-;

 R^{56} and R^{57} are independently C_1 - C_4 alkyl group optionally substituted with at least one halogen atom, C_2 - C_6 alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl optionally substituted with at least one halogen atom or phenyl group optionally substituted with at least one halogen atom, hydrogen atom, C_3 - C_6 cycloalkyl group, -XR⁵⁰ group or -NR⁶¹R⁶² group;

 R^{58} is hydrogen atom, C_1-C_6 alkyl group, C_2-C_6 alkenyl group, C_3-C_6 alkynyl group, C_1-C_4 alkylcarbonyl group, cyano- C_1-C_3 alkyl group, C_1-C_4 alkoxycarbonyl- C_1-C_4 alkyl group, $di-C_1-C_4$ alkoxycarbonyl- C_1-C_4 alkyl group, benzyl group, C_1-C_4 alkoxy- C_1-C_4 alkynyl group, $-(CH_2)_a-R^{75}$ group, $-(CH_2)_a-X^2-R^{72}$ group, $-(CH_2)_a-X^2-(CH_2)_b-R^{72}$ group or $-(CH_2)_a-X^2-(CH_2)_b-X^2-(CH_3)_b-X^2-(CH_3)_b-X^2-(CH_3)_c-R^{72}$ group;

 $$R^{59}$$ is hydrogen atom, $C_1\text{-}C_4$ alkyl group, $C_2\text{-}C_6$ alkenyl group, $C_3\text{-}C_6$ alkynyl group, cyano- $C_1\text{-}C_3$ alkyl group, $C_7\text{-}C_4$ alkylcarbonyl- $C_1\text{-}C_3$ alkyl group or phenyl group;

 $$\rm R^{60}$$ is $\rm C_1\text{--}C_4$$ alkyl group optionally substituted with at least one halogen atom;

 R^{61} and R^{62} are, the same or different, hydrogen atom or C,-C, alkyl group;

 R^{63} is C_1 - C_4 alkyl group optionally substituted

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with at least one halogen atom, C_1-C_4 alkoxy- C_1-C_4 alkyl group, C_1-C_4 alkylthio- C_1-C_4 alkyl group, C_3-C_6 cycloalkyl group, phenyl group whose ring may be substituted with one substituent selected from the group consisting of halogen atom, nitro group, cyano group, C_1-C_4 alkyl group, C_1-C_4 alkoxy group and halo- C_1-C_4 alkyl group, $-NR_1^{13}R_1^{74}$ group or - $(CH_4)_{-1}(O)_4-R_1^{75}$ group;

 $$R^{64}$$ is $C_1 - C_4$$ alkoxycarbonyl group or carboxyl group;

 R^{65} is chloromethyl group, cyanomethyl group, C_3 - C_6 cycloalkyl group into which at least one oxygen atom may be inserted, or C.-C. alkoxycarbonyl-C,-C, alkyl group;

 R^{66} is hydroxyl group or $-NR^{67}R^{68}$ group;

A is $-NR^{67}R^{68}$ group or $-S(0)_f-R^{69}$ group;

 R^{67} and R^{68} are, the same or different, hydrogen atom or $C_1\!-\!C_4$ alkyl group;

 R^{69} is C_1-C_4 alkyl group or C_1-C_4 haloalkyl group;

 R^{70} is hydrogen atom, hydroxyl group, halogen atom, C_1-C_4 alkyl group optionally substituted with at least one C_1-C_4 alkoxy group, C_3-C_6 cycloalkyl group into which at least one oxygen atom may be inserted, C_3-C_6 cycloalkyl group optionally substituted with one or two methyl groups, furyl group, thienyl group or $-C(=0)R^{71}$ group;

 $$R^{71}$$ and $$R^{72}$$ are, the same or different, $C_1\!-\!C_4$ alkyl group or $C_1\!-\!C_4$ alkoxy group;

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 $$R^{73}$$ and $$R^{74}$$ are, the same or different, $C_1\text{-}C_4$ alkyl group or phenyl group;

 R^{75} is C_3-C_6 cycloalkyl into which at least one oxygen atom may be inserted, C_3-C_6 cycloalkyl group optionally substituted with one or two methyl groups, furyl group, thienyl group or -C (=0) R^{71} group;

R76 is C1-C4 alkyl group;

a, b and c is independently 1, 2 or 3;

d is 0 or 1;

e is 2 or 3;

f is 1 or 2; and

 ${\rm X}^{\rm 2}$ is oxygen atom or sulfur atom.

In addition, as other N-substituted pyrazoles, there are the 3-substituted-2-aryl-4,5,6,7-tetrahydro-indazoles described in Lyga et al., Pesticide Sci., 42: p 29 (1994), and the like.

As specific examples of the compounds inhibiting electron transfer in photochemical system I, for example, there are paraquat, diquat, and the like. As specific examples of the compounds inhibiting electron transfer in photochemical system II, for example, there are triazine compounds (e.g., atrazine, etc.), urea compounds (e.g., diuron, etc.), nitrile compounds (e.g., bromoxynil and ioxynil) and the like. As specific examples of the compounds inhibiting 4-HPPD, for example, there are

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isoxazoles (e.g., isoxaflutole), pyrazoles, triketones, and

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the like. As specific examples of the compounds inhibiting PDS, for example, there are norflurazon, flurochloridone, fluridone, flurtamone, diflufenican, and the like. specific examples of the compounds inhibiting EPSPS, for example, there are glyphosate, and the like. As specific examples of the compounds inhibiting ALS, for example, sulfonylureas, imidazolinones, there are pyrimidinylthiobenzoates, triazolopyrimidines, and the like. As specific examples of the compounds inhibiting GS, for example, there are bialaphos, glufosinate, and the like. As specific examples of the compounds inhibiting DHP, for example, there are asulam, and the like. As specific examples of the compounds inhibiting ACC, for example, there are cyclohexanediones, aryloxyphenoxypropionates, and the like. As specific examples of the compounds inhibiting cellulose, for example, there are dichlobenil, and the like. Various examples of the weed control compounds

Various examples of the weed control compounds useful in the present invention are shown by the following chemical structures:

Structure 2

Structure 3

Structure 4

Structure 5

Structure 6

Structure 10

Structure 12

Structure 14

Structure 9

Structure 11

Structure 13

Structure 18

Structure 20

Structure 22

Structure 17

Structure 19

Structure 21

Structure 26

Structure 28

Structure 30

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Structure 27

Structure 29

Structure 34

Structure 36

Structure 33

Structure 35

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In the first aspect of the method of the present invention, the genes to be used are those encoding proteins having the following characteristics (a) to (c) (hereinafter sometimes referred to as the objective proteins):

- (a) having a specific affinity for weed control substances;
- (b) having substantially no capability of modifying substances for which said protein has a specific affinity; and
- (c) being substantially free from framework regions of variable regions of an immunoglobulin.

The term "a specific affinity" for weed control substances of the above characteristic (a) means that an enzyme (the objective protein) and a substrate (the weed control substance), or an enzyme (the objective protein) and an inhibitor or a regulator of an activity of the enzyme (the weed control substance) bind to each other, enzymatically; or that the objective protein and the weed control substance bind to each other on the basis of affinity and specificity, such as those shown in a receptor-chemical bond, for example, a bond between a receptor and a ligand, and the like. The objective proteins may be naturally occurring proteins; variants thereof obtained by introduction of amino acid substitution,

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addition, deletion, modification and the like into naturally occurring proteins; and artificially synthesized proteins having random amino acid sequences selected with the guidance of their affinity for weed control substances, in so far as they have structures specifically binding to weed control substances.

The term "having substantially no capability of modifying" in the characteristic (b) means that enzymatic reactivity with substances for which said protein has a specific affinity is substantially inactive or not existed (except the specific affinity for weed control substances in the characteristic (a)). Examples of this include a case that the objective protein does not have capability of converting a substance for which said protein has a specific affinity such as a certain weed control substance or a substance having an essential part of the structure of the substrates on the basis of a specific affinity for said protein, and the like to a substance having a chemical structure different from that of the substance for which said protein has a specific affinity. protein "having substantially no capability of modifying" can be, for example, identified by checking nonrecovery of the growth of a microorganism whose gene encoding the said protein is deleted and thus cannot grow under a usual condition in a case where the gene encoding

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the said protein is introduced into the microorganism in such a state that the introduced gene is expressed in the microorganism.

The term "substantially free from the framework regions of variable regions of an immunoglobulin" in the characteristic (c) mean that the objective protein does not form a stereostructure specific for the variable regions of an immunoglobulin. The term "framework regions of variable regions of an immunoglobulin" mean regions remaining after removing the hypervariable regions from the variable regions of H chain and L chain which are the constituents of the immunoglobulin molecule. In these regions, conservation of the amino acid sequences is relatively high and these regions function for maintaining the highly conserved stereostructure of the variable regions. formation of the above stereostructure, the hypervariable regions separately located at three sites on respective H chain and L chain are collected to one site on the stereostructure to form an antigen binding site [Alberts, B., et al. ed. (1983), Molecular Biology of the Cell, p 979, Garland Publishing, Inc., New Yorkl.

The objective protein having the above characteristic (c) can be selected on the basis of, for example, the amino acid sequences of the proteins. As specific examples of the protein, there are a protein which

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does not contain any amino acid sequence composed of about 30 amino acids or more and having about 60% or more homology with the known amino acid sequences of framework regions of the variable regions of immunoglobulin, and the like. For example, the presence or absence of the above framework regions can be confirmed by PCR using a gene encoding the protein as a template and DNAs having nucleotide sequences encoding the variable derived from H chain or L chain regions of immunoglobulin as amplification primers, for example, the primers VH1BACK and VH1FOR-2, or VK2BACK and VK4FOR described by Clackson, T. et al., Nature 352; p 624 (1991), or primers contained in a commercially available kit for cloning recombinant antibody genes, for example, Heavy primer mix or Light primer mix of Recombinant Phage Antibody System (Pharmacia Biotech) to analyze presence or absence of amplification of DNA having a given length. Examples of the binding proteins having a specific affinity for weed control substances also include peptides having an affinity for the weed control substances.

Specific examples of the objective proteins having the above characteristics of (a) to (c) include inactive-type binding proteins having an affinity for protoporphyrin IX [e.g., inactive-type magnesium chelatase whose substrate is protoporphyrin IX (the weed control

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substance), inactive-type ferrochelatase (protoheme ferrolyase; EC 4.9.9.1), inactive-type cobalt chelatase which catalyzes a chelating reaction of a cobalt ion with a compound having tetrapyrrole ring as a substrate, peptides having an affinity for protoporphyrin IX, i.e., proteins composed of 4 to 100 amino acids (for example, peptide HASYS having an affinity for protoporphyrin IX, e.g., a protein comprising the amino acid sequence of SEQ ID NO: 53 and a protein having the amino acid sequence of SEQ ID NO: 54; peptide RASSL having an affinity for protoporphyrin IX, e.g., a protein comprising the amino acid sequence of SEQ ID NO: 55 and a protein having the amino acid sequence of SEQ ID NO: 56; peptide YAGY having an affinity for porphyrin compounds, e.g., a protein comprising the amino acid sequence of SEQ ID NO: 57 and a protein having the amino acid sequence of SEQ ID NO: 58; peptide YAGF having affinity for porphyrin compounds, e.g., а protein comprising the amino acid sequence of SEQ ID NO: 59 and a protein having the amino acid sequence of SEQ ID NO: 60; and the like)], inactive-type binding proteins having an affinity for protoporphyrinogen IX (e.g., inactive-type PPO, inactive-type coproporphyrinogen III oxidase), and the like. The above inactive-type binding proteins include

The above inactive-type binding proteins include variants thereof whose activities have been lost by amino acid substitution, addition, deletion, modification and the

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like of naturally occurring active proteins under natural or artificial conditions.

Cellular dysfunction caused by weed control substances can be prevented by binding of these binding proteins to the weed control substances in plant cells to exhibit the desired weed control compound-resistance.

The inactive-type magnesium chelatase is protoporphyrin IX binding subunit protein of magnesium chelatase, or its variant having a specific affinity for protoporphyrin IX, and specific examples thereof include the subunit protein from which its organelle transit signal sequence has been deleted, and the like.

The inactive-type ferrochelatase is its variant having no capability of modifying protoporphyrin IX and having a specific affinity for protoporphyrin IX, and specific examples thereof include a ferrochelatase variant in which a region presumed to be a Fe ion binding site of ferrochelatse has been modified, and the like.

The inactive-type cobalt chelatase is a substrate binding subunit protein of cobalt chelatase, or its variant having no capability of modifying protoporphyrin IX and having a specific affinity for protoporphyrin IX.

The inactive-type PPO is its variant having no capability of oxidizing protoporphyrinogen IX and having a specific affinity for protoporphyrinogen IX, and specific

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examples thereof include a PPO variant in which a region presumed to be FAD binding site of PPO (a region having the amino acid sequence GXGXXG wherein X is any amino acid, e.g., a region comprising the 63rd to 68th amino acids from the N-terminus of chloroplast localized PPO of mouse-ear cress (Arabidopsis thaliana) and having the amino acid sequence of GGGISG) has been deleted, and the like.

The inactive-type coproporphyrinogen III oxidase is its variant having no capability of oxidizing protoporphyrinogen IX and having a specific affinity for protoporphyrinogen IX.

The genes encoding the above proteins can be obtained by, for example, as follows.

As the genes encoding protoporphyrin IX binding subunit protein of magnesium chelatase, for example, those derived from the photosynthetic bacterium, Rhodobacter capsulatus (Genebank accession M74001), mouse-ear cress (Genebank accession Z68495), barley (Genebank accession U96216), snapdragon (Antirrhinum majus) (Genebank accession U26916), Synechocystis P.C.C. 6803 (Genebank accession U29131) and the like have been known. For isolating such a known gene (its nucleotide sequence has been known), PCR can be carried out by using genomic DNA or cDNA of an organism having the desired gene as a template and primers produced on the basis of nucleotide sequences corresponding

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to those about the N- and C-termini of the protein encoded by the gene to amplify the desired gene. Further, genes encoding protoporphyrin IX binding subunit protein of magnesium chelatase can be obtained from photosynthetic organisms other than the above. For example, first, a cDNA library is constructed by obtaining mRNA from the desired photosynthetic organism, synthesizing cDNA by using the mRNA as a template with a reverse transcriptase, integrating the cDNA into a phage vector such as ZAPII, etc. or a plasmid vector such as pUC, etc. For amplifying a DNA fragment containing at least a part of the gene encoding protoporphyrin IX binding subunit protein of magnesium chelatase, PCR can be carried out by using the aboveconstructed cDNA library as a template and primers designed and synthesized on the basis of nucleotide sequences well conserved among known genes such as the above-described genes. Screening of the cDNA library can be carried out by using the DNA fragment thus obtained as a probe to select positive clones. The desired gene of protoporphyrin IX binding subunit protein of magnesium chelatase can be confirmed by sequence determination of the nucleotide sequence of the selected clone.

For obtaining the gene encoding a variant of protoporhyrin IX binding subunit protein of magnesium chelatase having an specific affinity for protoporphyrin IX,

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for example, the gene encoding the subunit protein is mutagenized by introduction of nucleotide substitution, addition, deletion, modification and the like, followed by introducing the resultant gene into Escherichia coli BL21(DE3) strain according to the method described by Gibson, L.C. D. et al., Proc. Natl. Acad. Sci. USA, 92; p 1941 (1995) and the like to obtain transformants, and culturing the transformants under conditions that high expression of the gene thus introduced occurs. The desired gene encoding a variant of the subunit protein having a specific affinity for protoporphyrin IX can be obtained by selecting a strain whose cultured cells have turned red and have the fluorescence absorption showing accumulation of protoporphyrin IX (excitation wavelength 405 nm, emission wavelength 630 nm).

As the genes encoding ferrochelatase, for example, those derived from Escherichia coli (Genebank accession D90259), Bacillus subtilis (Genebank accession M97208), Bradyrhizobium japonicum (Genebank accession M92427), yeast Saccharomyces cerevisiae (Genebank accession J05395), mouse (Genebank accession J05097), human being (Genebank accession D00726), barley (Genebank accession D26105), cucumber (Genebank accession D26106), and the like have been known. For isolating such a known gene (its nucleotide sequence has been known), PCR can be carried out

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by using genomic DNA or cDNA of an organism having the desired gene as a template and primers produced on the basis of nucleotide sequences corresponding to those about the N- and C-termini of the protein encoded by the gene to amplify the desired gene. Further, for obtaining other genes encoding ferrochelatase, for example, first, a cDNA library is constructed by obtaining mRNA from the desired organism, synthesizing cDNA by using the mRNA as a template with a reverse transcriptase, and integrating the cDNA into a phage vector such as ZAPII, etc. or a plasmid vector such as pUC, etc. The cDNA library can be introduced into ferrochelatase deficient mutant strain of Escherichia coli VS200 described by Miyamoto, K, et al., Plant Physiol., 105; p 769 (1994), followed by subjecting a complementation test to select clones containing ferrochelatase gene derived from the desired organism. Further, for amplifying a DNA fragment, PCR can be carried out by using the aboveconstructed cDNA library as a template and primers prepared on the basis of nucleotide sequences well conserved among known genes such as the above-described genes. Screening of the cDNA library can be carried out by using the DNA fragment thus obtained as a probe to select positive clones. The desired ferrochelatase gene can be confirmed by sequence determination of the nucleotide sequence of the selected clone.

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For obtaining the gene encoding a variant of ferrochelatase having no capability of modifying protoporphyrin IX and having a specific affinity for protoporphyrin IX (for example, the gene encoding a ferrochelatase variant in which the region presumed to be a Fe ion binding site of ferrochelatase is modified), PCR can be carried out by preparing a mutagenesis primer for introduction of mutation into the region on the basis of nucleotide sequence encoding the amino acid sequence about the region, and using a commercially available sitedirected mutagenesis kit (Mutan-Super Express, Shuzo) to obtain the gene encoding the above variant. Specifically, a wild type ferrochelatase gene is inserted into the cloning site of plasmid vector pKF19K and PCR is carried out by using the resultant plasmid DNA as template, the above-described mutagenesis primer and a selection primer for restoration of amber mutation located on kanamycin resistant gene of pKF19K. The gene amplified by PCR is introduced into Escherichia coli MV1184 (suppressor free strain) and the transformants are screened according to kanamycin resistance to isolate Escherichia coli having ferrochelatase gene in which the nucleotide sequence corresponding to the amino acid sequence which constitutes the desired region has been modified. isolated gene can be confirmed as the gene encoding the

desired protein by analyzing the nucleotide sequence of the plasmid DNA of the Escherichia coli.

The genes encoding the peptides having an affinity for protoporphyrin IX, i.e., the proteins composed of 4 to 100 amino acids can be obtained by synthesizing a peptide library according to, for example, the combinatorial chemistry method as described by Sugimoto, N., Nakano, S., Chem., Lett., p 939 (1997) and the like, selecting a peptide having an affinity for the weed control substance, analyzing the amino acid sequence of the peptide thus selected with a peptide sequencer, designing a gene containing a nucleotide sequence encoding the amino acid sequence, and synthesizing the nucleotide sequence with a DNA synthesizer or the like.

Further, a phase clone displaying a peptide having an affinity for the weed control substance can be selected from a phage library according to phage display method. Specifically, for example, a phage library displaying a protein having a random amino acid sequence on the surface of M13 phage particles is constructed by inserting a nucleotide sequence encoding the protein having the random amino acid sequence into the upstream from the region encoding the coat protein pIII of M13 phage gene. On the other hand, the weed control substance labeled with biotin is bound to a plate coated with avidin or

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streptoavidin to prepare a support coated with the weed control substance. A phage displaying the desired protein having an affinity for the weed control substance can be isolated by screening the above phage library on the plate coated with the weed control substance and the gene of the desired protein can be obtained from the isolated phage.

The gene encoding a protein containing the repetition of the amino acid sequence represented by SEO ID NO: 53, 55, 57 or 59 four times or eight times can be produced by, for example, selecting a nucleotide sequence in which the nucleotide sequence encoding the above amino acid sequence is repeated the given times after the initiation codon ATG, synthesizing an oligonucleotide comprising the selected nucleotide sequence oligonucleotide comprising а nucleotide sequence complementary to the selected nucleotide sequence by a DNA synthesizer, and then subjecting them to annealing. Further, the genes encoding the amino acid sequence represented by SEQ ID NO: 54, 56, 58 or 60 can be produced by selecting a nucleotide sequence encoding the amino acid sequence, synthesizing an oligonucleotide comprising the selected nucleotide sequence and another oligonucleotide comprising a nucleotide sequence complementary to the selected nucleotide sequence by a DNA synthesizer, and then subjecting them to annealing. In this respect, for

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selecting the nucleotide sequence encoding the given amino acid sequence, for example, it is preferred to select codons frequently used in genes derived from plants.

As PPO genes, for example, those derived from Escherichia coli (Genebank accession X68660), subtilis (Genebank accession M97208), Haemophilus influenzae (Genebank accession L42023), mouse (Genebank accession D45185), human being (Genebank accession D38537), mouse-ear cress (Genebank accession D83139), tobacco (Genebank accession Y13465, Y13466) and the like have been For isolating such a known gene (its nucleotide known. sequence has been known), PCR is carried out by using genomic DNA or cDNA of an organism having the desired gene a template and primers produced on the basis of nucleotide sequences corresponding to those about the N- $\,$ and C-termini of the protein encoded by the gene to amplify the desired gene. Further, for obtaining other PPO genes, for example, first, a cDNA library is constructed from an organism having the desired gene according to the abovedescribed method. The cDNA library can be introduced into Escherichia coli PPO deficient mutant strain described by Narita, S., et al., Gene, 182; p 169 (1996), followed by subjecting a complementation test to select clones containing PPO gene derived from the desired organism. Further, for amplifying a DNA fragment, PCR can

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be carried out by using the above-constructed cDNA library as a template and primers prepared on the basis of nucleotide sequences well conserved among known genes such as the above-described genes. Screening of the cDNA library can be carried out by using the DNA fragment thus obtained as a probe to select positive clones. The desired PPO gene can be confirmed by sequence determination of the nucleotide sequence of the selected clone.

For obtaining the gene encoding a variant of PPO having no capability of oxidizing protoporphyrinogen IX and having a specific affinity for protoporphyrinogen IX, for example, PPO gene is mutagenized by introducing nucleotide substitution, addition, deletion, modification, etc. and the resultant modified gene is introduced into the above Escherichia coli whose growth is inhibited dependently by treatment with a PPO inhibitory-type herbicidal compound. A gene encoding a protein having protoporphyrinogen IX binding capability can be selected by culturing the Escherichia coli thus obtained in presence of hemin, aminolevulinic acid and inhibitory-type herbicidal compound to select a clone which can grow even in the light. A gene encoding a protein having no capability of oxidizing protoporphyrinogen IX can be selected by expressing the modified gene thus selected in a host such as Escherichia coli, etc. to prepare a

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protein encoded by the gene, and measuring its capability of oxidizing protoporphyrinogen IX according to the method described by Jacobs, N.J. and Jacobs, J.M. (1982) Enzyme, 28, 206-219 and the like. More specifically, the above modified gene is inserted into an expression vector for Escherichia coli and introduced into PPO gene (hemG locus) deficient mutant of Escherichia coli such as Escherichia coli BT3 strain described by Yamamoto, F., et al., Japanese J. Genet., 63; p 237 (1988) and the like. The Escherichia coli is cultured in a culture medium containing hemin and aminolevulinic acid in addition to the cell growth inhibitor corresponding to the selection marker of the vector introduced into the Escherichia coli to obtain transformants. The protein encoded by the modified gene can be produced from the transformant. Further, a gene which does not complement PPO gene deficiency of its host cell can be obtained by culturing the transformant in a culture medium substantially free from hemin and aminolevulinic acid to identify a strain which does not grow. This latter method can also be used for selection of the gene encoding a protein having no capability of oxidizing protoporphyrinogen IX.

Further, for obtaining the gene encoding a variant of PPO in which the region presumed to be a FAD binding site of PPO (the region having the amino acid

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sequence GXGXXG, wherein X is any amino acid) is deleted, first, a mutagenesis primer for introduction of deletion mutation of the region is prepared on the basis of the nucleotide sequence encoding the amino acid sequence about the region. Then, PCR is carried out by using the mutagenesis primer and a commercially available site-directed mutagenesis kit (Mutan-Super Express, Takara Shuzo) as described above to obtain the gene encoding the above variant protein in which the region has been deleted.

The genes encoding peptide proteins such as the peptides HASYS and RASSL having an affinity for protoporphyrin IX, and the peptides YAGA and YAGF having an affinity for prophyrin compounds, and the like can be obtained by subjecting oligonucleotides synthesized by a DNA synthesizer to annealing.

Furthermore, genes encoding unknown peptide proteins having affinities for other weed control substances can be produced by the following methods and the like. For example, various peptide libraries can be constructed according to, for example, the combinatorial chemistry method as described by Sugimoto, N., Nakano, S., Chem., Lett., p 939 (1997), and the like. Peptides are selected from the peptide libraries thus constructed with the guidance of affinities for weed control substances, followed by analyzing the amino acid sequences of the

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peptides with a peptide sequencer. Thus, genes encoding the peptides can be synthesized by a DNA synthesizer. Alternatively, phase clones displaying peptides having affinities for weed control substances can be obtained by selecting phage libraries according to phage display method. Specifically, for example, a phage library displaying a protein having a random amino acid sequence on the surface of M13 phage particles is constructed by inserting a nucleotide sequence encoding the protein having the random amino acid sequence into the upstream from the region encoding the coat protein pIII of M13 phage gene. other hand, a weed control substance labeled with biotin is bound to a plate coated with avidin or streptoavidin to prepare a support coated with the weed control substance. A phage displaying the desired protein having an affinity for the weed control substance can be isolated by screening the above phage library on the plate coated with the weed control substance and the gene of the desired protein can be obtained from the isolated phage.

As the genes encoding coproporphyrinogen III oxidase, for example, those derived from Escherichia coli (Genebank accession X75413), Salmonella typhimurium (Genebank accession L19503), yeast Saccharomyces cerevisiae (Genebank accession J03873), mouse (Genebank accession D16333), human being (Genebank accession D16333), soybean

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(Genebank accession X71083), barley (Genebank accession X82830), tobacco (Genebank accession X82831) and the like have been known. For isolating such a known gene (its nucleotide sequence has been known), PCR is carried out by using genomic DNA or cDNA of an organism having the desired gene as a template and primers produced on the basis of nucleotide sequences corresponding to those about the Nand C-termini of the protein encoded by the gene to amplify desired gene. Further, for obtaining the coproporphyrinogen III oxidase genes, for example, first, a cDNA library is constructed from an organism having the desired gene by preparing mRNA from the desired organism, synthesizing cDNA using the mRNA as a template with a reverse transcriptase and integrating this into a plasmid vector such as pRS313 described by Sikorski, R.S., et al., Genetics, 122; p 19 (1989), and the like. The cDNA library can be introduced into yeast coproporphyrinogen III oxidase deficient mutant strain HEM13 described by Troup, B., et al., Bacteriol., 176; p 673 (1994), followed by subjecting complementation test to select clones containing coproporphyrinogen III oxidase derived from the desired organism. Further, for amplifying a DNA fragment, PCR can be carried out by using the above-constructed cDNA library a template and primers prepared on the basis of nucleotide sequences well conserved among known genes such

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as the above-described genes. Screening of the cDNA library can be carried out by using the DNA fragment thus obtained as a probe to select positive clones. The desired coproporphyrinogen III oxidase gene can be confirmed by sequence determination of the nucleotide sequence of the selected clone.

For obtaining the gene encoding a variant of coporphyrinogen III oxidase having no capability oxidizing protoporphyrinogen IX and having a specific affinity for protoporphyrinogen IX. for coproporphyrinogen III oxidase gene is mutagenized by introducing nucleotide substitution, addition, deletion, modification, etc. and the resultant gene is introduced into the above Escherichia coli whose growth is inhibited light-dependently by treatment with a PPO inhibitory-type herbicidal compound. A gene encoding a protein having protoporphyrinogen IX binding capability can be selected by culturing the Escherichia coli thus obtained in the hemin, aminolevulinic acid and presence of PPO inhibitory-type herbicide to select a clone which can grow even in the light. A gene encoding a protein having no capability of oxidizing protoporphyrinogen IX can be selected by expressing the modified gene thus selected in a host such as Escherichia coli, etc. to prepare a protein encoded by the gene, and measuring its capability of

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oxidizing protoporphyrinogen IX according to the method described by Jacobs, N.J. and Jacobs, J.M. (1982) Enzyme, 28, 206-219 and the like.

The genes which is used in the second aspect of the method of the present invention are those encoding proteins having the following characteristics (a) to (c):

- (a) having a specific affinity for protoporphyrinIX:
- (b) having substantially no capability of modifying protoporphyrinogen IX; and
- (c) being substantially free from framework regions of variable regions of immunoglobulins.

The term "a specific affinity" for protoporphyrin IX in the characteristic (a) is substantially the same as that in the above first aspect of the method of the present invention and means that the protein and protoporphyrin IX bind to each other, enzymatically or the protein and protoporphyrin IX bind to each other on the basis of affinity and specificity as those shown in receptor chemical bond such as a bond between a receptor and a ligand and the like. The proteins may be naturally occurring proteins; variants thereof in which amino acid substitution, addition, deletion, modification and the like are introduced into naturally occurring proteins; and artificially synthesized proteins having random amino acid

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sequences which are selected with the guidance of an affinity for protoporphyrin IX in so far as they have structures specifically binding to protoporphyrin IX.

The term "having substantially no capability of modifying" protoporphyrinogen IX in the characteristic (b) means that enzymatic reactivity with protoporphyrinogen IX of the protein is substantially inactive or not existed. For example, this means that the protein does not have capability of converting protoporphyrinogen IX into a substance having a chemical structure different from that of protoporphyrinogen IX.

The term "substantially free from framework regions of variable regions of immunoglobulins" means the same as that in the above first aspect of the method of the present invention and the protein does not form the stereostructure specific for the variable regions in the immunoglobulin as is described hereinabove.

As specific examples of the proteins having the above characteristics (a) to (c), there are active or inactive-type binding proteins having an affinity for protoporphyrin IX [e.g., active or inactive-type magnesium chelatase whose substrate is protoporphyrin IX, active or inactive-type ferrochelatase, active or inactive-type cobalt chelatase which catalyzes a chelating reaction of a cobalt ion with a compound having tetrapyrrole ring as a

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substrate, peptides, i.e., proteins composed of 4 to 100 amino acids, having an affinity for protoporphyrin IX (for example, proteins containing at least one peptide selected from peptide HASYS having an affinity for protoporphyrin IX, e.g., a protein comprising the amino acid sequence of SEQ ID NO: 53 and a protein having the amino acid sequence of SEO ID NO: 54; peptide RASSL having an affinity for protoporphyrin IX, i.e., a protein comprising the amino acid sequence of SEQ ID NO: 55 and a protein having the amino acid sequence of SEQ ID NO: 56; peptide YAGY having an affinity for porphyrin compounds, e.g., a protein comprising the amino acid sequence of SEQ ID NO: 57 and a protein having the amino acid sequence of SEQ ID NO: 58; peptide YAGF having affinity for porphyrin compounds, i.e., a protein comprising the amino acid sequence of SEQ ID NO: 59 and a protein having the amino acid sequence of SEQ ID NO: 60; and the like)], and the like.

The genes encoding the above proteins can be obtained by, for example, as follows.

Active-type magnesium chelatases are composed of three heterogenous subunit proteins, i.e., protoporhyrin IX binding subunit protein (H subunit protein), I subunit protein and D subunit protein, all of them are essential for catalytic acitivity. Three independent subunit proteins are encoded by different genes. The genes of

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protoporphyrin IX binding subunit protein can be obtained by PCR or screening of cDNA library as described hereinabove.

As the gene encoding I subunit protein of a magnesium chelatase, for example, those derived from photosynthetic bacterium, Rhodobacter sphaeroides (Genebank accession AF017642), Rhodobacter capsulatus accession Z11165), Arabidopsis (Genebank accession D49426), barley (Genebank accession U26545), soybean (Genebank accession D45857), tobacco (Genebank accession AF14053), Synechocystis P.C.C.6803 (Genebank accession U35144) and the like have been known. For isoltaing such a known gene (its nucleotide sequence has been known), PCR can be carried out by using genomic DNA or cDNA of an organism having the desired gene as a template and primers produced on the basis of nucleotide sequences corresponding to those about the N- and C-termini of the protein encoded by the desired gene. Further, genes encoding I subunit protein of a magnesium chelatase can be obtained from photosynthetic organisms other than the above. For example, first, a cDNA library is constructed by obtaining mRNA from the desired photosynthetic organisms, synthesizing cDNA by using the mRNA as a template with a reverse transcriptase, integrating the cDNA into a phage vector such as ZAPII, etc. or plasmid vector such as pUC, etc. For amplifying a DNA

fragment containing at least a part of the gene encoding I subunit protein of a magnesium chelatase, PCR can be carried out by using the above-constructed cDNA library as a template and primers designed and synthesized on the basis of nucleotide sequences well conserved among known genes such as the above described genes. Screening of the cDNA library can be carried out by using the DNA fragment thus obtained as a probe to select positive clones. The desired gene of I subunit protein of a magnesium chelatase can be confirmed by determination of the nucleotide sequence of the selected clone.

As the gene encoding D subunit protein of a magnesium chelatase, for example, those derived from photosynthetic bacterium, Rhodobacter sphaeroides (Genebank accession AJ001690), Rhodobacter capsulatus (Geneband accession Z11165), pea (Genebank accession AFO14399), (Genebank accession Y10022), Synechocystis tobacco P.C.C.6803 (Genebank accession X96599) and the like have The isolation of such a known gene (its been known. nucleotide sequence has been known) or genes other than the above can be carried out in the same manner as described in that of the gene encoding I subunit protein of magnesium chelatase.

The genes used in the third aspect of the method of the present invention are those encoding proteins having

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the following characteristics (a) to (c):

- (a) having a specific affinity for protoporphyrinogen IX;
- (b) having the capability of modifying coproporphyrinogen III; and
 - (c) being substantially free from framework regions of variable regions of immunoglobulins.

affinity" term "a specific The in the characteristic (a) is protoporphyrinogen IX substantially the same as that in the above first or second aspect of the method of the present invention and means that the protein and protoporphyrinogen IX bind to each other, enzymatically or the protein and protoporphyrinogen IX are bound to each other on the basis of affinity and specificity as those shown in receptor-chemical bond such as a bond between a receptor and a ligand and the like. The proteins may be naturally occurring proteins; variants thereof in which amino acid substitution, addition, deletion, modification and the like are introduced into naturally occurring proteins; and artificially synthesized proteins having random amino acid sequences which are quidance of an affinity selected with the protoporphyrinogen IX in so far as they have structures specifically binding to protoporphyrinogen IX.

The term "having the capability of modifying"

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coproporphyrinogen III in the characteristic (b) means that enzymatical reactivity with coproporphyrinogen III of the proteins is active. For example, this means that the protein has the capability of converting coproporphyrinogen III into a substance having a chemical structure different from that of coproporphyrinogen III.

The term "substantially free from framework regions of variable regions of immunoglobulins" means the same as that in the above first or second aspect of the method of the present invention and the protein does not form the stereostructure specific for the variable regions in the immunoglobulin as is described hereinabove.

As specific examples of the proteins having the above characteristics (a) to (c), there are active or inactive-type binding proteins having an affinity for proporphyrinogen IX, for example, active-type coproporphyrinogen III oxidase whose substrate is proporphyrinogen IX, and the like.

As a reference, the activity of a magnesium chelatase, a ferrochelatase or a coproporphyrinogen III oxidase is, for example, measured by using the following method.

(1) A magnesium chelatase:

The genes encoding independent three subunit proteins are used to detect a magnesium chelatase activity

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according to the method by Gibson, L.C.D., et al. (Proc. Natl. Acad. Sci. USA, 92; p 1941 (1995)) and the like.

(2) A ferrochelatse:

A ferrochelatase activity can, for example, be detected according to the method by Porra, R.J. (Anal. Biochem., 68; p 289 (1975)) and the like.

(3) A coproporphyrinogen III oxidase:

A coproporphyrinogen III oxidase activity can, for example, be detected according to the method by Yoshinaga, T., Sano, S., et al. (J. Biol. Chem., 255; p 4722 (1980)) and the like.

In the fourth aspect of the method of the present invention, there may be used in addition to the gene encoding the protein having the characteristics (a) to (c) (as described in the first to third aspects of the present invention), at least one altered form of an enzymatic activity selected from an altered PPO activity, an altered EPSPS activity and an altered glyphosate oxidoreductase (GOX) activity. Said altered form of the enzymatic activity in the plant cell can give a resistance to a weed control compound in an amount inhibiting a naturally occurring form of said enzymatic activity. Typically, such an amount of the weed control compound is an amount which can set forth a herbicidal control over the growth of a plant cell, which by inhibiting a naturally occurring form

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of the enzymatic activity. In this regard, to give the resistance to a PPO inhibitory-type herbicidal compound, it is preferable that the plant cell additionally comprises the altered PPO activity. Likewise, to give the resistance to glyphosate, it is preferable that the plant cell additionally comprises the altered EPSPS activity or the altered GOX activity.

Glyphosate is the common name given to the weed control compound, N-(phosponomethyl)glycine. In this regard, glyphosate includes the ammonium salt, sodium salt, isopropylamine salt, trimethylsulfonium salt, potassium salt or the like salt. Further, glyphosate is a compound encompassed by said compound inhibiting EPSPS, as described above.

The term "altered form of an enzymatic activity" means that the enzymatic activity is different from that which naturally occurs in a plant cell, which altered form of an enzymatic activity provides a resistance to a weed control compound that inhibits the naturally occurring activity thereof. Said naturally occurring enzymatic activity in the plant cell is the enzymatic activity which occurs naturally in the absence of direct or indirect manipulation by man of such naturally occurring enzymatic activity.

A second gene in the plant cell is useful to

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confer said altered form of the enzymatic activity therein. As such, the second gene typically provides in the plant cell, a gene providing for the altered PPO activity, for the altered EPSPs activity or for the altered GOX activity. Various proteins can be encoded by the second gene, so that the second gene can provide for said altered form of enzymatic activity when expressed in the plant cell.

In utilizing the second gene for the altered PPO activity, the second gene can encode a naturally occurring protein substantially having PPO activity. In the plant cell, such a protein substantially having PPO activity can be a protein having a capability of oxidizing protoporphyrinogen IX and which has a specific affinity for protoporphyrinogen IX.

In its amino acid sequence, the protein substantially having PPO activity preferably contains said region presumed to be FAD binding site of PPO. Such a protein substantially having PPO activity may be PPO. As genes encoding PPO, there can be utilized the known "PPO genes" as described above. Further, there can be utilized a naturally occurring protein substantially having PPO activity which activity is resistant to the PPO inhibitory-type herbicidal compound (as described in EP 0770682 or WO 9833927)

Further, the protein substantially having PPO

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activity can have substituted, deleted or added thereto amino acids, such that the resulting variant protein has substantially the PPO activity. Conventional methods well known in the art can be used to substitute, delete or add the amino acids thereto. USP 5939602 and WO 9704089 describe variant PPO substantially having PPO activity which activity is uninhibited by a PPO inhibitory-type herbicidal compound. The second gene may encode such a variant PPO.

In utilizing the second gene for the altered EPSPS activity, the second gene can encode a naturally occurring protein substantially having EPSPS activity. Such a protein substantially having EPSPS activity is a protein having a capability to modify in the plant cell, phosphoenolpyruvic acid (PEP) with 3-phosphoshikimic acid to 5-enolpyruvyl-3-phosphoshikimic acid.

Such a protein substantially having EPSPS activity may be EPSPS. As a polynucleotide encoding EPSPS, there can be utilized a known polynucleotide encoding EPSPS. Examples of such a polynucleotide encoding EPSPS include those derived from Petunia hybrida (Genebank accession M37029), Mitchell diploid petunia (as described in EP218571), Salmonella typhymurium (as described in EP508909), Tomato (strain VF36) pistil (Genebank accession M21071), Arabidopsis thaliana (Genebank accession X06613),

soy beans, Zea mays (Genebank accession X63374),

Escherichia coli (Genebank accession X00557), Agrobacterium tumefaciens sp.strain CP4 (class II) and the like.

Additionally, the second gene can also encode a naturally occurring protein substantially having EPSPS activity which activity is resistant to glyphosate, such as a bacterial EPSPS which activity is resistant to glyphosate.

Further, the second gene when providing for the altered EPSPS activity in the plant cell can also encode a variant protein substantially having EPSPS activity. As such, the protein substantially having EPSPS activity can have substituted, deleted or added thereto amino acids such that the resulting protein substantially has the EPSPS activity. Examples of such a variant protein substantially having EPSPS activity include a variant EPSPS which activity is resistant to glyphosate, a variant EPSPS in which a chloroplast transit peptide is added thereto and the like.

The variant EPSPS which activity is resistant to glyphosate can be produced by substituting, deleting or adding nucleotides to a gene encoding EPSPS. For example, a substitutive mutation can be introduced to a polynucleotide encoding EPSPS to produce a variant polynucleotide. The protein encoded by the resulting variant gene can then be confirmed for a resistance to

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glyphosate and for the EPSPS activity.

The resistance to glyphosate may be confirmed by introducing the variant gene to a particular Escherichía coli mutant and by culturing the resulting particular Escherichia coli mutant in a specified minimal nutrient MOPS medium which has glyphosate added thereto. As the particular variant Esherichia coli in this case, there is utilized an Escherichia coli mutant which is deficient in its endogenous EPSPS gene (aroA locus) and which has the growth thereof inhibited in the specified minimal nutrient MOPS medium (in which there is no glyphosate therein). Further, in this case, the specified minimal nutrient MOPS medium is specified in that there is no aromatic amino acids present therein. When glyphosate is added to the minimal nutrient MOPS medium, the glyphosate is in an amount which would typically inhibit in normal growing conditions, the growth of an Escherichia coli mutant which is deficient in its endogenous EPSPS gene but which has introduced thereto a naturally occurring gene encoding herbicidally sensitive EPSPS. By selecting the resulting clones growable in such specified minimal nutrient MOPS medium containing glyphosate, there can be obtained a variant polynucleotide encoding a variant EPSPS having an activity which is resistant to glyphosate. The EPSPS activity can be confirmed by introducing said variant gene

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to a host cell and then by according to the method described in EP 409815. In this regard, there can be obtained the second gene encoding the variant EPSPS substantially having EPSPS activity which activity is resistant to glyphosate.

In utilizing the second gene for the altered GOX activity, the second gene can encode a naturally occurring protein substantially having GOX activity. Such a protein substantially having the GOX activity is a protein having a capability to degrade glyphosate to less herbicidal products, such as aminomethyl phosphonate (AMPA) and glyoxylate. In cases in which glyphosate is degraded into AMPA and glyoxylate, for example, the protein substantially having GOX activity may cleave the C-N bond of glyphosate.

Such a protein substantially having GOX activity may be GOX. As a polynucleotide encoding GOX, there can be utilized a known polynucleotide encoding GOX. Examples of naturally occurring GOX genes include those derived from Pseudomonas sp.strains LBAA, Pseudomonas sp.strains LBr, Agrobacterium sp.strain T10 and the like.

Further, the second gene when providing for the altered GOX activity in the plant cell can encode a variant protein substantially having GOX activity. As such, the protein substantially having GOX activity can have

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substituted, deleted or added thereto amino acids such that the resulting protein substantially has the GOX activity.

As an example of such a variant protein substantially having GOX activity, there is mentioned a variant GOX in which a chloroplast transit peptide is added thereto.

Conventional methods well known in the art can be used to substitute, delete or add the amino acids thereto.

The GOX activity of a protein substantially having GOX activity can be confirmed by introducing a gene encoding the protein substantially having GOX activity into a specified Escherichia coli mutant and by culturing the resulting specified Escherichia coli mutant in a minimal nutrient MOPS medium containing glyphosate as the sole nitrogenous source therein. As the specified Escherichia coli mutant in this case, there is utilized an Escherichia coli mutant which can grow in a minimal nutrient MOPS medium having a non-herbicidal aminophosphate compound as the sole nitrogenous source therein, such as E.coli SR2000 Mpu+ . The glyphosate therein is in an amount which would typically inhibit the growth of the specified Escherichia coli mutant having no said gene encoding the protein substantially having GOX activity introduced thereto. By selecting the resulting clones growable in such minimal nutrient MOPS medium containing glyphosate as the sole nitrogenous source therein, there can be obtained a

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polynucleotide encoding a protein substantially having GOX activity. Such results suggest that in cases in which the protein substantially having GOX activity degrades glyphosate into AMPA, the growable clones use to grow, AMPA as a nitrogenous source. In practice, a 3-14C labeled glyphosate may be used to confirm that said growable clone consumes and degrades glyphosate. For example, the growable clone may be cultured with the 3-14C labeled glyphosate and the cell extract thereof may then be analyzed with HPLC.

The second gene encoding an above protein substantially having PPO activity, EPSPS activity or GOX activity can be obtained, for example, as follows.

For isolating a known gene encoding a protein substantially having PPO activity, EPSPS activity or GOX activity, PCR can be carried out by using genomic DNA or cDNA of an organism having the desired gene as a template and primers produced on the basis of nucleotide sequences corresponding to those about the N- and C-termini of the protein to amplify the desired gene. Further, genes encoding a protein substantially having PPO activity, EPSPS activity or GOX activity can be obtained from organisms other than the above. For example, first, a cDNA library is constructed by obtaining mRNA from an organism and synthesizing cDNA by using the mRNA as template with

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reverse transcriptase and integrating the cDNA into a phage vector such as ZAP II, etc. or a plasmid vector such as pUC, etc. For the protein substantially having PPO activity, the cDNA library may be introduced into Escherichia coli PPO deficient mutant strain VSR800 described by Narita, S., et al., Gene, 182; p 169 (1996), followed by subjecting a complementation test to select clones containing PPO gene derived from the desired organism. Further, for amplifying a DNA fragment containing at least a part of the desired gene, PCR can be carried out by using the above-constructed cDNA library as a template and primers designed and synthesized on the basis of nucleotide sequences well conserved among known genes such as the above-described genes. Screening of the cDNA library can be carried out by using the DNA fragment thus obtained as a probe to select positive clones. The desired gene, i.e., a gene encoding the protein substantially having the PPO activity, EPSPS activity or GOX activity, can be confirmed by determination of the nucleotide sequence of the selected clone.

Examples of methods used to confer the altered EPSPS activity or altered GOX activity include the following. An example may include a method of introducing into a cultivated plant a gene having a polynucleotide sequence encoding a petunia (Mitchell diploid petunia)

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EPSPS downstream of a high expression promoter such as a cauliflower mosaic virus 35S promoter (EP 218571). A further example may include a method of introducing into a cultivated plant a gene having a 35S promoter upstream of a polynucleotide sequence encoding an Agrobacterium (Agrobacterium tumefaciens sp.strain CP4) EPSPS fused with a chloroplast transit peptide of a petunia (Petunia hybrida) EPSPS (WO 9204449, USP 5633435). A furthermore example may include a method of introducing into a cultivated plant a gene having 2 continuous 35S promoters upstream a polynucleotide encoding a sunflower chloroplast transit peptide of small subunit of ribulose-1,5bisphosphate carboxylase (ssRUBISCO), the 22 amino acids from the N-terminus of maize ssRUBISCO, maize chloroplast transit peptide of ssRUBISCO and a Salmonella (Salmonella typhyrium) EPSPS (EP 508909). Even furthermore, an example may include a method of introducing into a cultivated plant, a gene having downstream from a promoter of Arabidopsis thaliana alcohol dehydrogenase A, a polynucleotide encoding an Arabidopsis thaliana chloroplast transit peptide and GOX (WO 9706269). Yet even furthermore, an example may include a method of introducing into a cultivated plant, the above gene encoding GOX as well as a gene having downstream from a 35S promoter possessing an enhanced promoter activity with the omega sequence of tobacco mosaic virus, a

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polynucleotide sequence which encodes an Agrobacterium (Agrobacterium tumefaciens sp. strain CP4) EPSPS (class II) downstream a chloroplast transit peptide of Petunia (Petunia hybrida) EPSPS (WO 9706269). Still yet even furthermore, an example may include a method of introducing into a cultivated plant, a gene encoding a variant EPSPS having amino acid substitutions therein which augment the resistant to glyphosate [Hinchee, M.A.W. et al., BIO/TECHNOLOGY, 6: p915 (1988), EP 389066, EP 409815, WO 9206201 and USP 5312910].

Examples of methods used to confer the altered PPO activity include the following. An example may include a method of over-expressing in a plant cell, a gene encoding wild-type, naturally occurring PPO (USP 5767373). A further example may include a method of expressing in a plant cell, a variant protein substantially having PPO activity which activity is not inhibited by a PPO inhibitory-type herbicidal compound (USP 5939602). A furthermore example may include a method of expressing a PPO substantially having PPO activity which is not inhibited by a PPO inhibitory-type herbicidal compound, wherein the PPO is derived from bacteria (EP 0770682 or WO 9833927).

In the method (including the above first to third aspects) of the present invention, for introducing the gene

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encoding the protein having the characteristics of (a) to (c) into a plant cell, a gene encoding one protein can be Further, plural genes encoding different introduced. proteins can be introduced into a plant cell. When said altered form of enzymatic activity is given to the plant cell, the second gene encoding one protein may also be introduced. Further, plural genes of the second gene can be introduced into the plant cell to provide for said altered form of enzymatic activity therein. In introducing the gene encoding the protein having the characteristics of (a) to (c) and second gene thereto, the gene encoding the protein having the characteristics of (a) to (c) may be introduced into the plant cell with the second gene, or may be introduced before or after the second gene is introduced to the plant cell. Such gene introduction into plant cells can be carried out by conventional gene engineering techniques, for example, Agrobacterium infection (JP-B 2-58917 and JP-A 60-70070), electroporation into protoplasts (JP-A 60-251887 and JP-A 5-68575), particle gun methods (JP-A 5-508316 and JP-A 63-258525), and the like.

Preferably, the gene to be introduced into a plant cell is integrated into a vector having a selection marker gene such as a gene which can give cell growth inhibitor resistance to the plant cell. For example, the gene encoding the protein having the characteristics of (a)

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to (c) and the second gene, when utilized for the altered form of enzymatic activity, can be integrated into one of such vectors. Further, the gene encoding the protein having the characteristics of (a) to (c) and the second gene may also each be integrated, respectively, into such vectors having a selection marker gene. In integrating the gene encoding the protein having the characteristics (a) to (c) and the second gene into such respective vectors, the selection marker gene utilized for the vector for the second gene is typically different from the selection marker gene utilized for the vector for the gene encoding the protein having the characteristics (a) to (c).

For expression of the gene encoding the protein having the characteristics (a) to (c) in the plant cell, the gene can be introduced into a chromosome of a plant cell by homologous recombination [Fraley, R.T. et al., Proc. Natl. Acad. Sci. USA, 80; p 4803 (1983)] to select the plant cell expressing the gene. Alternatively, the gene can be introduced into a plant cell in the form that it is operably ligated to a promoter and a terminator both of which can function in the plant cell.

The term "operably ligated" used herein means that the above promoter and terminator are joined in such a state that the introduced gene is expressed in the plant cell under control of the promoter and the terminator.

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To provide for the altered form of enzymatic activity, the second gene is expressed in a plant cell. For expression of the second gene in the plant cell, the second gene can likewise be introduced into a chromosome of a plant cell by homologous recombination to select the plant cell expressing the second gene. Alternatively, the second gene can be introduced into a plant cell in the form that it is operably ligated to a promoter and a terminator both of which can function in the plant cell. utilized, the second gene is typically expressed at a level such that the amount of the protein encoded by the second gene provides for the altered form of enzymatic activity and further confer the resistance of the plant cell. It is preferable when the second gene encodes PPO or EPSPS, that the second gene provide for the altered form of enzymatic activity through over-expression. If so desired, transcriptionally strong promoter which can function in the plant cell can be utilized with the second gene.

As the promoter which can function in a plant cell, for example, there are constitutive promoters derived from T-DNA such as nopaline synthase gene promoter, octopine synthase gene promoter, etc., promoters derived from plant viruses such as 19S and 35S promoters derived from cauliflower mosaic virus, etc., inductive promoters such as phenylalanine ammonia-lyase gene promoter, chalcone

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synthase gene promoter, pathogenesis-related protein gene promoter, etc., and the like. The promoter is not limited these promoters and other plant promoters can be used.

As the terminator which can function in a plant cell, for example, there are terminators derived from T-DNA such as nopaline synthase terminator, etc., terminators derived from plant viruses such as terminators derived from garlic viruses GV1, GV2, etc., and the like. The terminator is not limited to these terminators and other plant terminators can be used.

As the plant cells into which the gene encoding the protein having the characteristics of (a) to (c) are introduced, for example, there are plant tissues, whole plants, cultured cells, seeds and the like. Examples of the plant species into which the genes are introduced include dicotyledones such as tobacco, cotton, rapeseed, sugar beet, mouse-ear cress, canola, flax, sunflower, potato, alfalfa, lettuce, banana, soybean, pea, legume, pine, poplar, apple, grape, citrus fruits, nuts, etc.; and monocotyledones such as corn, rice, wheat, barley, rye, oat, sorghum, sugar cane, lawn, etc. The second gene may also be introduced into such plant cells.

The transformant plant cells expressing the gene encoding the protein having the characteristics of (a) to (c) can be obtained by culturing cells into which the gene

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is transferred in a selection culture medium corresponding to a selection marker joined to the locus on the gene, for example, a culture medium containing a cell inhibitor, or the like, and isolating a clone capable of growing in the culture medium. Further, the selection culture medium should also correspond to a selection marker joined to the locus of the second gene when the altered form of enzymatic activity is also present transformant plant cells. Alternatively, the transformant plant cells can be selected by culturing plant cells into which the gene is introduced in a culture medium weed control compound to which containing the and isolating clones capable resistance is given, growing in the culture medium.

The desired weed control compound-resistant plant can be obtained from the transformant cells thus obtained by regenerating the whole plant according to a conventional plant cell culture method, for example, that described in Plant Gene Manipulation Manual, Method for Producing Transgenic Plants, UCHIMIYA, Kodansha Scientific (1996). Thus, the transformed plants such as plant tissues, whole plants, cultured cells, seeds and the like can be obtained.

For example, rice and mouse-ear cress expressing the gene encoding the protein having the characteristics of (a) to (c) can be obtained according to the method

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described Experimental Protocol of Model Plants, Rice and Mouse-Ear Cress Edition, (Supervisors: Koh SHIMAMOTO and Kiyotaka OKADA, Shujun-sha, 1996), Chapter 4. according to the method described in JP-A 3-291501, soybean expressing the gene encoding the binding protein by introducing the gene into soybean adventitious embryo with a particle gun. Likewise, according to the method described by Fromm, M.E., et al., Bio/Technology, 8; p 838 (1990), corn expressing the gene encoding the above protein can be obtained by introducing the gene into adventitious embryo with a particle gun. Wheat expressing the gene encoding the above protein can be obtained by introducing the gene into sterile-cultured wheat immature scutellum with a particle gun according to a conventional method described by TAKUMI et al., Journal of Breeding Society (1995), 44: Extra Vol. 1, p 57. Likewise, according to a conventional method described by HAGIO, et al., Journal of Breeding Society (1995), 44; Extra Vol. 1, p 67, barley expressing the gene encoding the above protein can be obtained by introducing the gene into sterile-cultured barley immature scutellum with a particle gun.

For confirmation of weed control compoundresistance of the plant expressing the gene encoding the above protein, preferably, the plant is reproduced with applying the weed control compound to which resistance is

given to evaluate the degree of reproduction of the plant. For more quantitative confirmation, for example, in case of resistance to a compound having PPO inhibitory-type herbicidal activity, preferably, pieces of leaves of the plant are dipped in aqueous solutions containing the compound having PPO inhibitory-type herbicidal activity at various concentrations, or the aqueous solutions containing the compound having herbicidal activity are sprayed on pieces of leaves of the plant, followed by allowing to stand on an agar medium in the light at room temperature. After several days, chlorophyll is extracted from the plant leaves according to the method described by Mackenney, G., J. Biol. Chem., 140; p 315 (1941) to determine the content of chlorophyll.

Since the weed control compound-resistant plants (e.g., plant tissues, whole plants, cultured cells, seeds, etc.) obtained by the method of the present invention (including the first to fourth aspects) show resistance to weed control compounds, even in case that a weed control compound is applied to a growth area (e.g., cultivation area, proliferation area, etc.), the plant can grow. Therefore, when a weed control compound is applied to a growth area of the desired weed control compound resistant-plant, the desired plant can be protected from plants without resistance to the weed control plant. For example,

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weeds can be controlled efficiently by applying a weed control compound on a growth area of the plant having resistance to the weed control compound.

Further, by applying a weed control compound to a growth area of the weed control compound-resistant plant obtained by the method of the present invention (including the first to third aspects) and other plants (e.g., those having no or weak resistance to the weed control compound), one of the plants can be selected on the basis of the difference in growth between the plants. For example, by applying (adding) a weed control compound to a cultivation area (culture medium) of the weed control compound-resistant plant cells obtained by the method of the present invention and other plant cells (e.g., those having no or weak resistance to the weed control compound), one of the plant cells can be selected efficiently on the basis of the difference in growth between the plants.

The following Examples further illustrate the present invention in detail but are not to be construed to limit the scope thereof.

Example 1

 $\label{eq:subunit} Isolation \mbox{ of Protoporphyrin IX Binding Subunit}$ Protein Gene of Magnesium Chelatase.

Genomic DNA of photosynthetic bacterium

Rhodobacter sphaeroides ATCC17023 was prepared using

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ISOPLANT kit for genomic DNA preparation (manufactured by Nippon Gene). Then, according to the description of Gibson, L.C.D. et al., Proc. Natl. Acad. Sci. USA, 92; p 1941 (1995), PCR was carried out by using about 1 µg of said template, and 10 pmol of genomic DNA as а oligonucleotide composed of nucleotide sequence represented by SEO ID NO: 1 and 10 pmol of an oligonucleotide composed of nucleotide sequence represented by SEQ ID NO: 2 as fragment containing to amplify the DNA primers protoporphyrin IX binding subunit protein gene bchH of magnesium chelatase. The oligonucleotides were prepared with a DNA synthesizer (PE Applied Biosystems: Model 394 DNA/RNA Synthesizer) and purified with an oligonucleotide purification cartridge (PE Applied Biosystems: Cartridge). The PCR was carried out by maintaining at 94°C for 2 minutes, at 96°C for 40 seconds and then at 68°C for 7 minutes, repeating a cycle for maintaining at 96°C for 40 seconds and then at 68°C for 7 minutes 28 times, and finally maintaining at 96°C for 40 seconds, at 68°C for 7 minutes and then at 72°C for 10 minutes.

Example 2

Expression of Protoporphyrin IX Binding Subunit Protein Gene of Magnesium Chelatase in $Escherichia\ Coli$ (hereinafter abbreviated to $E.\ coli)$

According to the description of Gibson, L.C.D. et

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al., Proc. Natl. Acad. Sci. USA, 92; p 1941 (1995), the DNA fragment containing bchH gene prepared in Example 1 was digested with the restriction enzymes NdeI and BglII. resultant DNA fragment was inserted between NdeI restriction site and BamHI restriction site of expression vector pET11a (manufactured by Stratagene) to obtain plasmid pETBCH (Fig. 1). This plasmid pETBCH introduced into E. coli BL21(DE3) strain competent cells (manufactured by Stratagene) according to the manual attached to the competent cells to obtain E. coli BL21(DE3)/pETBCH strain. The strain was inoculated into 1.5 ml LB liquid culture medium (1% tryptone, 0.5% yeast extract, 0.5% NaCl) containing 100 µg/ml ampicillin in a tube (14 x 10 mm), and the tube was covered with aluminum foil (hereinafter referred to as dark conditions), cultured with shaking at 37°C under light of fluorescent lamp (about 8000 When the absorbance at 600 nm of the liquid culture medium became about 0.6, isopropyl β-D-thiogalactopyranoside (IPTG) was added to the liquid culture medium so that the final concentration was 0.4 mM, and the culture continued for about additional 20 hours. At that time, the Escherichia coli turned red and fluorescent absorbance (excitation wavelength 405 nm, emission wavelength 630 nm) which showed the accumulation of protoporphyrin IX in E. coli was observed. When E. coli BL21(DE3)/pETBCH strain

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was cultured according to the same manner except that IPTG was not added, *E. coli* did not turned red and the above fluorescent absorbance did not detected. In contrast to this, when *E. coli* BL21(DE3)/pETBCH strain was cultured according to the same manner (with IPTG) except that the tube was not covered with aluminum foil (hereinafter referred to as light conditions), *E. coli* grew and turned red as above.

Example 3

Sovbeans (Glycine max var. Williams82) were seeded and cultivated at 25°C for 20 days and green leaves were collected. The collected green leaves were frozen with liquid nitrogen and the frozen leaves were ground in a mortar with a pestle. From the ground leaves, RNA were extracted by using RNA extracting reagent ISOGEN (manufactured by Nippon Gene) according to the manual attached thereto. The resultant RNA liquid extract was subjected to ethanol precipitation to collect total RNA, then the total RNA was fractionated by using poly (A) RNA fractionating kit BIOMAG mRNA Purification Kit (manufactured by Perceptive Bio System) according to the manual attached thereto to collect poly (A) RNA fraction. Using 1 µg of this poly (A) RNA fraction as a template, cDNA was

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synthesized with the cDNA synthetic reagent contained in Marathon cDNA amplification kit (manufactured by Clontech) according to the manual attached thereto. PCR was carried out by using the resultant cDNA as a template, and an oligonucleotide composed of nucleotide sequence of SEQ ID NO: 3 and an oligonucleotide composed of nucleotide sequence of SEO ID NO: 4 as primers to amplify the DNA fragment containing chloroplast-type protoporphyrinogen IX The above oligonucleotides were prepared oxidase gene. with a DNA synthesizer (PE Applied Biosystems: Model 394 DNA/RNA Synthesizer) and purified with an oligonucleotide purification cartridge (PE Applied Biosystems: OPC Cartridge). The PCR was carried out by maintaining at 94°C for 1 minutes and then at 65°C for 5 minutes, repeating a cycle for maintaining at 94°C for 15 seconds and then at 65°C for 5 minutes 29 times. After the PCR, the amplified DNA fragment was purified by filtering the reaction mixture with MicroSpin S-400HR (manufactured by Pharmacia Biotech), and the DNA fragment was ligated to plasmid pCR2.1 (manufactured by Invitrogen) cleaved by restriction enzyme SalI to obtain plasmid pSPPO-P. Then, the plasmid was introduced into competent cells of $E.\ coli\ INV \alpha F'$ strain (manufactured by Invitrogen) and ampicillin resistant strains were selected. Then, the plasmid contained in selected ampicillin resistant strains was sequenced by using

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Dye terminator cycle sequencing kit (manufactured by PE Applied Biosystems) and DNA sequencer 373S (manufactured by PE Applied Biosystems). As a result, the nucleotide sequence of SEQ ID NO: 5 was revealed, thereby confirming that plasmid pSPPO-P contained chloroplast-type protoporphyrinogen IX oxidase gene of soybean.

The plasmid pSPPO-P was digested with restriction enzyme PshBI, the resultant DNA fragment was blunted by using T4 DNA polymerase and further digested with SphI to isolate the DNA fragment containing chloroplast-type PPO gene of soybean and lac promoter. Then, the plasmid pACYC184 (manufactured by Nippon Gene) was digested with the restriction enzymes NruI and SphI to remove a fragment of 410 bp and the above DNA fragment was inserted instead to obtain plasmid pACYCSP (Fig. 2). Then, the plasmid pACYCSP was introduced into PPO gene (hemG gene locus) deficient mutant E. coli BT3 strain (described in Yamamoto, F. et al., Japanese J. Genet., 63; p 237 (1988) etc.) according to the method described in Hanahan, D.J., Mol. Biol., 166; p 557 (1983). The resultant E. coli were cultured in YPT medium (5 g/liter yeast extract, 5 g/liter tryptone, 5 g/liter peptone, 10 g/liter NaCl, pH 7.0) containing 15 µg/ml chloramphenicol and 10 µg/ml kanamycin select E, coli BT3/pACYCSP strain resistant †o chloramphenicol and kanamycin whose hemG gene deficiency

was complemented by PPO gene derived from soybean.

Example 4

Test of Protoporphyrin IX Binding Subunit Protein of Magnesium Chelatase for Capability of Giving Weed Control Compound-Resistance

E. coli BT3/pACYCSP strain prepared in Example 3 was inoculated into YPT medium containing 10 or 1 ppm of PPO inhibitory type herbicidal compound represented by the above Structure 8, 10 µg/ml hemin, 50 µg/ml aminolevulinic acid, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin, dark conditions or cultured under light conditions according to the same manner as in Example 2. As a control, E. coli BT3/pACYCSP strain was cultured in the same medium as above without the herbicidal compounds under the same conditions. Then, 18 hours after initiation of culture, the absorbance of the liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 1.

Table 1

E. coli strain	Culture conditions	Relative absorbance			
		Concentration of test compound			
		10 ppm	1 ppm	0 ppm	
BT3/pACYCSP	in the light	0.10	0.25	1.0	
BT3/pACYCSP	in the dark	0.73	0.95	1.0	

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Plasmid pTVBCH (Fig. 3) was constructed by amplification of the DNA fragment containing both gene derived from photosynthetic bacterium Rhodobacter sphaeroides using the oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 1 and the oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 2 according to the same manner as in Example 1, digestion of the resultant DNA fragment with the restriction enzymes NcoI and BglII and introducing the digested DNA fragment between NcoI restriction site and BamHI restriction site of plasmid pTV118N (manufactured by Takara Shuzo Co., Ltd.).

Plasmids pTVBCH and pTV118N respectively were introduced into *E. coli* BT3/pACYCSP strain prepared in Example 3 according to the method described in Hanahan, D.J., Mol. Biol., 166; p 557 (1983). The resultant *E. coli* were cultured in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin to obtain *E. coli* BT3/pACYCSP+pTVBCH strain bearing plasmids pACYCSP and pTVBCH, and *E. coli* BT3/pACYCSP+pTV118N strain bearing plasmids pACYCSP and pTV118N.

These strains were inoculated into YPT medium containing 10 or 1 ppm of the PPO inhibitory-type herbicidal compound represented by the above Structure 8, 100 μ g/ml ampicillin, 15 μ g/ml chloramphenicol, 10 μ g/ml

kanamycin, 10 µg/ml hemin and 50 µg/ml aminolevulinic acid, cultured under dark conditions or light conditions according to the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of the liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 2.

Table 2

E. coli strain		Relative absorbance		
	Culture conditions	Concentration of test compound		
		10 ppm	1 ppm	0 ppm
BT3/pACYCSP+pTVBCH	in the light	0.80	0.77	1.0
BT3/pACYCSP+pTVBCH	in the dark	0.90	1.06	1.0
BT3/pACYCSP+pTV118N	in the light	0.18	0.31	1.0
BT3/pACYCSP+pTV118N	in the dark	0.68	0.77	1.0

Further, these strains were inoculated into YPT medium containing PPO inhibitory-type herbicidal compounds represented by the above Structures 1, 14, 15, 18-22, 29, 32, 33, 34 and 36, respectively, 100 µg/ml ampicillin, 15 µg/ml chloramphenicol, 10 µg/ml kanamycin, 10 µg/ml hemin and 50 µg/ml aminolevulinic acid, cultured under dark conditions or light conditions similar to the Example 2. Then, 18 hours after initiation of culture, the absorbance

of liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated.

5 The results are shown in Table 3.

Table 3

Test	m t	Relative absorbance				
compound	Test	BT3/		BT3/		
Structure	tructure concent-		pACYCSP+pTVBCH		pACYCSP+pTV118N	
No.	ration	in the	in the	in the	in the	
		light	dark	light	dark	
Structure 1	5.0	0.88	0.88	0.31	0.87	
Structure 14	10	0.47	0.93	0.12	0.81	
Structure 15	0.5	0.94	0.94	0.38	0.82	
Structure 18	2.0	0.68	1.0	0.33	0.91	
Structure 19	5.0	0.78	0.89	0.40	0.71	
Structure 20	5.0	0.57	0.88	0.11	0.75	
Structure 21	10	0.88	0.91	0.25	0.85	
Structure 22	10	0.55	0.93	0.29	0.94	
Structure 29	0.5	0.64	0.90	0.22	0.77	
Structure 32	2.0	0.70	0.94	0.37	0.87	
Structure 33	2.0	0.81	0.92	0.41	0.91	
Structure 34	1.0	0.41	0.94	0.19	0.86	
Structure 36	0.5	0.55	0.95	0.28	0.96	

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Example 5

Introduction of Gene Encoding Protoporphyrin IX
Binding Subunit Protein of Magnesium Chelatase into Tobacco

A plasmid was constructed for introducing bchH gene into a plant by Agrobacterium infection method. First, binary vector pBI121 (manufactured by Clontech) digested with restriction enzyme SacI, and KpnI linker (manufactured by Takara Shuzo Co., Ltd.) was inserted to prepare plasmid pBIK wherein SacI recognition site of pBI121 was removed and KpnI recognition site was added. other hand, according to the same manner as described in Example 1, PCR was carried out by using the genomic DNA of photosynthetic bacterium Rhodobacter sphaeroides as template, and the oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 7 and the oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 8 to amplify the DNA fragment containing bchH gene. the above plasmid pBIK was digested with restriction enzymes XbaI and KpnI to remove \$-qlucuronidase gene, and instead thereof, a DNA fragment which was obtained by digesting the above DNA fragment containing bchH gene with restriction enzymes XbaI and KpnI was inserted to produce plasmid pBIBCH (Fig. 4) in which bchH gene was joined downstream from 35S promoter. Binary vector pBI121 also digested with (manufactured by Clontech) was

restriction enzymes BamHI and SacI to remove β -glucuronidase gene, the resultant DNA fragment was blunted by using T4 DNA polymerase, followed by self-cyclization with T4 DNA ligase to construct plasmid pNO (Fig. 5). The plasmid was used as a vector control of bchH expression plasmid pBIBCH.

The plasmid pBIBCH and pNO were introduced into Agrobacterium tumefaciens LBA4404, respectively. Abrobacterium strain bearing pBIBCH and that bearing pNO were isolated by culturing the resultant transformants in a medium containing 300 µg/ml streptomycin, 100 µg/ml rifampicin and 25 µg/ml kanamycin and selecting the desired transformants.

Then, according to the method described in Manual for Gene Manipulation of Plant (by Hirofumi UCHIMIYA, Kodan-sha Scientific, 1992), the gene was introduced into tobacco. Agrobacterium strain bearing plasmid pBIBCH was cultured at 28°C overnight in LB medium and then leaf pieces of tobacco cultured sterilely were dipped in the liquid culture medium. The leaf pieces were cultured at room temperature for 2 days in Murashige-Skoog medium (MS-medium, described in Murasige T. and Skoog F., Physiol. Plant. (1962) 15, p 473) containing 0.8% agar, 0.1 mg/liter naphthalene acetic acid and 1.0 mg/liter benzyl aminopurine. Then, the leaf pieces were washed with sterilized water and

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cultured for 7 days on MS medium containing 0.8% agar, 0.1 mg/liter naphthalene acetic acid, 1.0 mg/liter benzyl aminopurine and 500 ug/ml cefotaxime. The leaf pieces were transplanted onto MS medium containing 0.8% agar, 0.1 mg/liter naphthalene acetic acid, 1.0 mg/liter benzyl aminopurine, 500 µg/ml cefotaxime and 100 µg/ml kanamycin (hereinafter referred to as selective MS medium) and cultured on the medium continuously for 4 months with transplanting the tobacco leaf pieces onto fresh selective MS medium every 1 month. During culture, stem-leaf differentiated shoots were appeared from the tobacco leaf pieces, these shoots were transplanted to MS medium containing 0.8% agar, 300 µg/ml cefotaxime and 50 µg/ml kanamycin to induce roots to obtain regenerated plants. resultant regenerated plant was transplanted and cultured on MS medium 0.8% agar and 50 µg/ml kanamycin to obtain tobacco plant into which bchH gene was introduced. Similarly, tobacco leaf pieces were infected with Agrobacterium strain bearing pNO to obtain regenerated plant from the tobacco leaf pieces and tobacco plant (hereinafter referred to as control recombinant tobacco).

Example 6

Test of Tobacco Bearing Introduced Gene Encoding Protoporphyrin IX Binding Subunit Protein of Magnesium Chelatase for Resistance to Herbicidal Compounds

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The tobacco leaves into which bchH gene was introduced and control recombinant tobacco leaves obtained in Example 5 were collected and each leaf was divided into the right and left equivalent pieces along the main vein, respectively. To one piece was applied an aqueous solution containing 0.3 ppm PPO inhibitory-type herbicidal compound of Structure 8, while, to the other piece was not applied the compound. These leaf pieces were placed on MS medium containing 0.8% agar and allowed to stand at room temperature for 7 days in light place. Then, each leaf piece was ground with pestle and mortar in 5 ml of 80% aqueous acetone solution to extract chlorophyll. extract liquid was diluted with 80% aqueous acetone solution and the absorbance was measured at 750 nm, 663nm and 645 nm to calculate total chlorophyll content according to the method described by Macknney G., J. Biol. Chem. (1941) 140, p 315. The results obtained from 4 clones of tobacco into which bchH gene was introduced (BCH1 to 4) and control recombinant tobacco is shown in Table 4. table, the resistant level to the herbicidal compound was represented by percentages of the total chlorophyll content of leaf pieces treated with herbicidal compound to that of untreated leaf pieces.

Table 4

Recombinant	Total chloro (mg/ g-fresh w	Resistant level to test	
tobacco	untreated- leaf	treated-leaf	compound(%)
control	2.49	0.19	7.63
BCH-1	1.35	1.70	126
BCH-2	2.06	2.14	104
BCH-3	1.93	1.57	81.3
BCH-4	1.51	1.06	70.2

The tobacco clone into which bchH gene was introduced and control recombinant tobacco were also treated in the same manner with the solution containing PPO inhibitory-type herbicidal compound represented by the above Structure 3, 7, 10, 11, 13, 17, 23, 24, 25, 27, 28, 30 or 35, and the resistant level to each herbicidal compound was measured. The results are shown in Table 5. In the table, the resistant levels to the herbicidal compound were represented by percentages of the total chlorophyll content of leaf pieces treated with the herbicidal compound to that of untreated leaf pieces.

Table 5

Test compound Structure No.	Test concentration	Resistant level to test compound (%)		
		bchH	control	
	(ppm)	recombinant	recombinant	
		tobacco	tobacco	
Structure 3	10	114	9.94	
Structure 7	30	89.3	8.62	
Structure 10	10	84.0	14.9	

Structure 11	0.30	78.1	5.51
Structure 13	30	95.2	14.8
Structure 17	0.30	80.4	14.3
Structure 23	3.0	106	5.58
Structure 24	10	129	5.18
Structure 25	10	104	16.0
Structure 27	10	86.8	16.8
Structure 28	0.30	72.2	8.79
Structure 30	3.0	102	4.24
Structure 35	0.30	83.3	17.4

Example 7

Isolation of Gene Encoding Variant Protein of Protoporphyrin IX Binding Subunit Protein of Tobacco Magnesium Chelatase

Total RNAs were prepared from leaf tissues of tobacco (Nicotiana tabacum cv. SR1) by using RNeasy Plant Kit (manufactured by QIAGEN) according to the manual attached thereto. The DNA fragment containing the gene encoding protoporphyrin IX binding subunit protein of tobacco magnesium chelatase whose chloroplast transit signal had been deleted (hereinafter referred to as the variant tobacco chelatase subunit) was obtained by using RNA LA PCR Kit (AMV) Ver 1.1 (manufactured by Takara Shuzo Co., Ltd.) according to the manual attached thereto. First, 1st strand cDNA was synthesized by using tobacco total RNAs as templates and Oligo dT-Adaptor Primer contained in the

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above kit as the primer with the reverse transcriptase contained in the above kit. Then, PCR was carried out by using the 1st strand cDNA as a template and LA Tag polymerase contained in the above kit to amplify the DNA fragment containing the gene encoding the variant tobacco chelatase subunit protein. In this PCR, oligonucleotide primer composed of the nucleotide sequence of SEO ID NO: 9 and the oligonucleotide primer composed of the nucleotide sequence of SEO ID NO: 10 were used. synthesized oligonucleotides were by using DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA Synthesizer) and purified with an oligonucleotide purification cartridge (PE Applied Biosystems; OPC cartridge). The PCR was carried out by maintaining at 94°C for 2 minutes and then repeating a cycle for maintaining at 94°C for 30 seconds, at 50°C for 30 seconds and then at 72°C for 7 minutes 30 times. After the PCR, the DNA fragment amplified by the PCR was cloned into plasmid pCR2.1 by using TA Cloning Kit (manufactured by Invitrogen) according to the manual attached thereto. The resultant plasmid was digested with restriction enzyme KpnI and analyzed by agarose gel electrophoresis. The plasmid from which 8.0 kb DNA fragment was detected was named pTCHLH. The plasmid had the structure that the gene encoding the variant tobacco chelatase subunit has been inserted in the direction

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expressible under the control of lac promoter. Plasmid pTCHLH was digested with restriction enzyme KpnI followed by self-ligaiton to obtain plasmid pTCHLH1 (Fig. 6) in which DNA fragment composed of about 60 nucleotides had been deleted from plasmid pTCHLH.

Example 8

Test of Variant Tobacco Magnesium Chelatase Subunit Protein for Capability of Giving Resistance to Herbicidal Compounds

The plasmid pTCHLH1 and pCR2.1 prepared in Example 7 were introduced into *E. coli* BT3/pACYCSP strain prepared in Example 3, respectively according to the method described in Hanahan, D.J., Mol. Biol., 166; p 557 (1983). *E. coli* BT3/pACYCSP+pTCHLH1 strain bearing plasmids pACYCSP and pTCHLH1, and *E. coli* BT3/pACYCSP+pCR2.1 strain bearing plasmids pACYCSP and pCR2.1 were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 50 µg/ml kanamycin, respectively.

These *E. coli* strains were inoculated into YPT medium containing 10 or 1 ppm of the PPO inhibitory-type herbicidal compound represented by Structure 8, 100 μ g/ml ampicillin, 15 μ g/ml chloramphenicol, 50 μ g/ml kanamycin, 10 μ g/ml hemin and 50 μ g/ml aminolevulinic acid, cultured under dark conditions or light conditions according to the

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same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of the liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 6.

Table 6

		Relative absorbance			
E. coli strain	Culture conditions	concent compoun	ration d	of test	
		10 ppm	1 ppm	0 ppm	
BT3/pACYCSP+pTCHLH1	in the light	0.69	0.89	1.0	
BT3/pACYCSP+pTCHLH1	in the dark	0.92	0.93	1.0	
BT3/pACYCSP+pCR2.1	in the light	0.03	0.08	1.0	
BT3/pACYCSP+pCR2.1	in the dark	1.0	1.0	1.0	

Example 9

Introduction of Gene Encoding Variant Tobacco
Magnesium Chelatase Subunit Protein into Tobacco

A plasmid for introducing the gene encoding a variant tobacco magnesium chelatase subunit protein into tobacco by Agrobacterium infection method was constructed. First, the DNA fragment containing the gene encoding the variant tobacco magnesium chelatase subunit protein was prepared by digesting plasmid pTCHLH1 prepared in Example 7 with restriction enzymes KpnI and SalI. On the other hand,

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binary vector pBI121 (manufactured by Clonetech) digested with restriction enzyme SmaI and KpnI linker (manufactured by Takara Shuzo Co., Ltd.) was inserted into this portion to prepare plasmid pBI121K in which SmaI recognition site of pBI121 was removed and KpnI recognition The plasmid pBI121K was digested with site was added. restriction enzyme SacI followed by blunting the DNA by adding nucleotides to the double-stranded DNA gap with DNA Then, the DNA was dephosphorylated with Polymerase I. alkaline phosphatase derived from calf intestine and cyclized by inserting phosphorylated SalI linker (4680P, manufactured by Takara Shuzo Co., Ltd.) to construct plasmid pBI121KS. The binary vector pBI121KS was digested with restriction enzymes KpnI and SalI to remove glucuronidase gene and the gene encoding the variant tobacco magnesium chelatase subunit protein was inserted into this portion to prepare plasmid pBITCHLH (Fig. 7).

The plasmid pBITCHLH was introduced into Agrobacterium tumefaciens LBA4404. The resultant transformants were cultured in a medium containing 300 µg/ml streptomycin, 100 µg/ml rifampicin and 25 µg/ml kanamycin, followed by selecting the desired transformants to isolate a Agrobacterium strain bearing pBITCHLH.

Leaf pieces of tobacco cultured sterilely are infected with the Agrobacterium strain and, according to

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the same manner as in Example 5, tobacco into which the gene encoding the variant tobacco magnesium chelatase subunit protein is introduced is obtained.

Example 10

Confirmation of Resistance to Herbicidal

Compounds of Tobacco Bearing Introduced Gene Encoding

Variant Tobacco Magnesium Chelatase Subunit Protein

The levels of resistance to herbicidal compounds are confirmed quantitatively by testing tobacco introduced with the gene encoding the variant tobacco magnesium chelatase subunit protein prepared in Example 9 according to the same manner as in Example 6.

Example 11

Isolation of Gene Encoding Variant Protein of Soybean PPO Having No Capability of Oxidizing Protoporphyrinogen IX and Having Specific Affinity for Protoporphyrinogen IX

PCR was carried out by using plasmid pSPPO-P prepared in Example 3 as a template and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 11 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 12 as primers to amplify the DNA fragment encoding soybean PPO whose chloroplast transit signal and FAD binding sequence had been deleted (hereinafter referred to as the variant soybean PPO). The oligonucleotides were

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prepared with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with oligonucleotide purification cartridge (PE Applied Biosystems; OPC cartridge). The PCR was carried out by repeating a cycle for maintaining at 94°C for 1 minute, at 55°C for 2 minutes and the 72°C for 3 minutes 30 times. amplified DNA fragments were digested with restriction Ncol and SalI, and introduced between NcoI restriction site and SalI restriction site of plasmid pTV118N (manufactured by Takara Shuzo Co., Ltd.) construct plasmid pTVGMP (Fig. 8).

The plasmid pTVGMP was introduced into $E.\ coli$ PPO gene deficient mutant BT3 strain according to the method described in Hanahan, D.J., Mol. Biol., 166; p 557 (1983). When the resultant $E.\ coli$ were cultured in YPT medium containing 100 µg/ml ampicillin and 10 µg/ml kanamycin, no growth complemented clone was obtained.

Example 12

Test for Effect of Giving Resistance to Herbicidal Compounds of Variant Soybean PPO

Plasmids pTVGMP and pTV118N prepared in Example 11 were introduced into E. coli BT3/pACYCSP strain prepared in Example 3 respectively according to the method described in Hanahan, D.J., Mol. Biol., 166; p 557 (1983). E. coli BT3/pACYCSP+pTVGMP strain bearing plasmids pACYCSP and

pTVGMP, and $E.\ coli$ BT3/pACYCSP+ pTV118N strain bearing plasmids pACYCSP and pTV118N were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin.

These E. coli strains were inoculated into YPT medium containing 10 or 1 ppm of PPO inhibitory-type herbicidal compound represented by Structure 8, 100 µg/ml ampicillin, 15 µg/ml chloramphenicol, 10 µg/ml kanamycin, 10 µg/ml hemin and 50 µg/ml aminolevulinic acid, cultured under dark conditions or light conditions according to the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 7.

Table 7

		Relative absorbance		
E. coli strain	Culture conditions	Concentration of test		
		10 ppm	1 ppm	,0 ppm
BT3/pACYCSP+pTVGMP	in the light	0.33	0.85	1.0
BT3/pACYCSP+pTVGMP	in the dark	0.91	0.94	1.0
BT3/pACYCSP+pTV118N	in the light	0.05	0.09	1.0
BT3/pACYCSP+pTV118N	in the dark	0.89	0.91	1.0

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Example 13

Introduction of the Gene Encoding Variant Soybean

A plasmid for introducing the gene encoding the variant soybean PPO into a plant by Agrobacterium infection method was constructed. PCR was carried out by using the plasmid pSPPO-P prepared in Example 3 as a template, an oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 13 and an oligonucleotide primer composed of the nucleotide sequence of SEO ID NO: 14 to amplify the DNA fragment containing the gene encoding the variant sovbean Then, plasmid pBI121K prepared in Example 9 was PPO. digested with the restriction enzymes KpnI and SacI to remove B-glucuronidase gene, and the DNA fragment which was obtained by digesting the DNA fragment containing the above gene encoding the variant soybean PPO with restriction enzymes KpnI and Sac I was inserted into this portion to prepare plasmid pBIGMP (Fig. 9) in which the gene was joined downstream from 35S promoter.

The plasmid pBIGMP was introduced into Agrobacterium tumefaciens LBA4404. The resultant transformants were cultured in a medium containing 300 µg/ml streptomycin, 100 µg/ml rifampicin and 25 µg/ml kanamycin, followed by selecting the desired transformants to isolate Agrobacterium strain bearing pBIGMP.

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Leaf pieces of tobacco cultured sterilely were infected with the Agrobacterium strain and, according to the same manner as in Example 5, tobacco into which the gene encoding the variant soybean PPO was introduced was obtained.

Example 14

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Gene Encoding
Variant Soybean PPO

The level of resistance to PPO inhibitory type herbicidal compound represented by Structure 8 was confirmed quantitatively by testing tobacco into which the gene encoding the variant soybean PPO prepared in Example 13 was introduced according to the same manner as in Example 6. The results obtained from 4 clones (GMP 1-4) of tobacco introduced with the gene encoding the variant soybean PPO and control recombinant tobacco are shown in Table 8. In the table, the resistant level to herbicidal compound is represented by percentage of the total chlorophyll content of leaf pieces treated with the herbicidal compound to that of untreated leaf pieces.

Table 8

Recombinant	Total chlorophyll content (mg/ g-fresh weight)			
tobacco	untreated- leaf treated-leaf		level to test compound(%)	
control	3.49	0.35	10.0	

2.0

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GMP-1	1.89	2.55	135
GMP-2	0.89	0.96	108
GMP-3	1.50	1.49	99.3
GMP-4	2.91	2.34	80.4

Example 15

Isolation of PPO Gene of Chlamydomonas

Chlamydomonas reinhardtii CC407 strain was obtained from Chlamydomonas Genetics Center (address: DCMB Group, Department of Botany, Box 91000, Duke University, Durham, USA), uE/m2/s 27708-1000, cultured under 200 photosynthesis active light for 5 days in TAP liquid culture medium (E. H. Harris, The Chlamydomonas Sourcebook, Academic Press, San Diego, 1989, p 576-577) containing 7 mM NH₄Cl, 0.4 mM MgSO₄·7H₂O, 0.34 mM CaCl₂·2H₂O, 25 mM potassium phosphate, 0.5 mM Tris (pH 7.5), 1 ml/liter Hatner miner element and 1 ml/liter glacial acetic acid to obtain 200 ml $(1.0 \times 10^6 \text{ cells/ml})$ liquid culture medium containing early stationary growth phase cells.

Total RNAs were prepared from these cells by using ISOGEN (manufactured by Nippon Gene) according to the manual attached thereto. Also, poly(A)RNA was fractionated using BioMag mRNA Purification Kit (manufactured by Perceptive Bio System) according to the manual attached thereto. cDNA was synthesized from the resultant poly(A)RNA by using Marathon cDNA Amplification Kit (manufactured by

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Clontech) according to the manual attached thereto and the cDNA was used as a template for PCR.

As PCR primers, an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 15 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 16 were prepared. The oligonucleotides were prepared with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotide purification cartridge (PE Applied Biosystems; OPC cartridge).

PCR was carried out by preparing a reaction liquid using Advantage cDNA PCR kit (manufactured by Clontech) according to the manual attached thereto, and then, after maintaining at 94°C for 1 minute and then at 65°C for 5 minutes, repeating a cycle for maintaining at 94°C for 15 seconds and the 65°C for 5 minutes 29 times. After the PCR, the amplified DNA fragments were purified by filtering the reaction liquid with MicroSpin S-400HR (manufactured by Pharmacia Biotech), and the DNA fragment was cloned into plasmid pCR2.1 by using TA Cloning Kit (manufactured by Invitrogen) according to the manual attached thereto to construct plasmid pCPPO.

The nucleotide sequence of DNA fragment contained in the resultant plasmid pCPPO was determined by using Dye terminator cycle sequencing kit (manufactured by PE applied

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Biosystems) and DNA sequencer 373S (manufactured by PE applied Biosystems). As a result, the nucleotide sequence of SEQ ID NO: 17 was revealed, thereby confirming that plasmid pCPPO contained the full length PPO cDNA of Chlamydomonas reinhardtii.

Example 16

Isolation of Gene Encoding Variant Protein of Chlamydomonas reinhardtii PPO Having No Capability of Oxidizing Protoporphyrinogen IX and Specific Affinity for Protoporphyrinogen IX

PCR was carried out by using plasmid pCPPO template, prepared in Example 15 as а oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 19 and an oligonucleotide composed of the nucleotide SEQ ID NO: 20 as primers to amplify the DNA fragment encoding Chlamydomonas reinhardtii PPO whose chloroplast transit signal and FAD binding sequence had been deleted (hereinafter referred to as the variant Chlamydomonas reinhardtii PPO). The oligonucleotides were prepared with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA purified with synthesizer) and an oligonucleotide Biosystems; purification cartridge (PE Applied cartridge). The PCR was carried out by repeating a cycle for maintaining at 94°C for 1 minute, at 55°C for 2 minutes and then at 72°C for 3 minutes 30 times. The amplified DNA

fragment was digested with restriction enzymes BamHI and SacI, and inserted between BamHI restriction site and SacI restriction site of plasmid pTV119N (manufactured by Takara Shuzo Co., Ltd.) to construct plasmid pTVCRP (Fig. 10).

The plasmid pTVCRP was introduced into $E.\ coli$ PPO gene deficient mutant BT3 strain according to the method described in Hanahan, D. J., Mol. Biol., 166; p 557 (1983). When the resultant $E.\ coli$ were cultured in YPT medium containing 100 µg/ml ampicillin and 10 µg/ml kanamycin, no growth complemented clone was obtained.

Example 17

Test of Variant Modified Chlamydomonas
reinhardti PPO for Capability of Giving Resistance to
Herbicidal Compounds

Plasmids pTVCRP and pTV118N prepared in Example 16 were introduced into E. coli BT3/pACYCSP strain prepared in Example 3 respectively according to the method described in Hanahan, D. J., Mol. Biol., 166; p 557 (1983). E. coli BT3/pACYCSP+pTVCRP strain bearing plasmids pACYCSP and pTVCRP, and E. coli BT3/pACYCSP+pTV118N strain bearing plasmids pACYCSP and pTV118N were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin.

These E. coli strains were inoculated into YPT medium containing 10 or 1 ppm of the PPO inhibitory-type

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herbicidal compound represented by Structure 8, 100 μ g/ml ampicillin, 15 μ g/ml chloramphenicol, 10 μ g/ml kanamycin, 10 μ g/ml hemin and 50 μ g/ml aminolevulinic acid, cultured under dark conditions or light conditions in the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of liquid culture medium was measured at 600 nm. By taking the absorbance of the medium containing no herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 9.

Table 9

		Relative absorbance			
E. coli strain	Culture conditions	Concentration of test			
		10 ppm	1 ppm	0 ppm	
BT3/pACYCSP+pTVCRP	in the light	0.23	0.42	1.0	
BT3/pACYCSP+pTVCRP	in the dark	0.81	0.82	1.0	
BT3/pACYCSP+pTV118N	in the light	0.12	0.24	1.0	
BT3/pACYCSP+pTV118N	in the dark	0.80	0.91	1.0	

Example 18

Introduction of Gene Encoding Variant
Chlamydomonas reinhardtii PPO into Tobacco

A plasmid for introducing the gene encoding the variant Chlamydomonas reinhardtii PPO into a plant by Agrobacterium infection method was constructed. The DNA

fragment containing the gene encoding the variant Chlamydomonas reinhardtii PPO was prepared by digesting plasmid pTVCRP prepared in Example 16 with restriction enzymes BamHI and SacI. Binary vector pBI121 (manufactured by Clontech) was digested with restriction enzymes BamHI and SacI to remove β -glucuronidase gene and the above gene encoding the variant Chlamydomonas reinhardtii PPO was inserted into this portion to prepare plasmid pBICRP (fig. 11).

The plasmid pBICRP was introduced into Agrobacterium tumefaciens LBA4404. The resultant transformants were cultured in a medium containing 300 µg/ml streptomycin, 100 µg/ml rifampicin and 25 µg/ml kanamycin, followed by selecting the desired transformants to isolate Agrobacterium strain bearing pBICRP.

Leaf pieces of tobacco cultured sterilely were infected with the Agrobacterium strain and, according to the same manner as in Example 5, tobacco into which the gene encoding the variant Chlamydomonas reinhardtii PPO was introduced was obtained.

Example 19

Confirmation of Resistance to Herbicidal

Compounds of Tobacco Bearing Introduced Gene Encoding

Variant Chlamydomonas reinhardtii PPO

The level of resistance to the PPO-inhibitory

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type herbicidal compound represented by Structure 8 was confirmed quantitatively by testing tobacco into which the gene encoding the variant Chlamydomonas reinhardtii PPO prepared in Example 18 was introduced according to the same manner as in Example 6. The results obtained from 4 clones (CRP 1-4) of tobacco into which the gene encoding the variant Chlamydomonas reinhardtii PPO was introduced and control recombinant tobacco is shown in Table 10. In the table, the resistant levels to the herbicidal compound are represented by percentages of the total chlorophyll content of leaf pieces treated with the herbicidal compound to that of untreated leaf pieces.

Table 10

Recombinant	Total chloron (mg/g-fresh w	Resistant level to test	
tobacco	untreated- leaf	treated-leaf	compound(%)
control	2.28	0.42	18.4
CRP-1	1.27	1.54	121
CRP-2	1.50	1.67	111
CRP-3	1.10	1.11	101
CRP-4	1.58	1.57	99.4

Example 20

Test of Variant Protein of Barley Ferrochelatase
Having Affinity for Protoporphyrin IX Specifically for
Capability of Giving Resistance to Herbicidal Compounds

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A plasmid bearing barley ferrochelatase gene was prepared by the method described in Miyamoto, K. et al., Plant Physiol. 105; p 769 (1994). The resultant plasmid was digested with restriction enzymes NspI and EcoRI to obtain the DNA fragment containing the gene encoding barley ferrochelatase whose signal sequence had been deleted (hereinafter referred tο as the variant. barlev This DNA fragment was inserted between ferrochelatase). SphI restriction site and EcoRI restriction site of plasmid pTV119N (manufactured by Takara Shuzo Co., Ltd.) to construct plasmid pTVHVF1 (Fig. 12).

The plasmids pTVHVF1 and pTV118N were introduced into E. coli BT3/pACYCSP strains prepared in Example 3 respectively according to the method described in Hanahan, D.J., Mol. Biol., 166; p 557 (1983). E. coli BT3/pACYCSP+pTVHVF1 strain bearing plasmid pACYCSP and pTVHVF1, and E. coli BT3/pACYCSP+pTV118N strain bearing plasmid pACYCSP and pTV118N were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin.

These *E. coli* strains were inoculated into YPT medium containing 10 or 1 ppm of the PPO inhibitory-type herbicidal compound represented by Structure 8, 100 μ g/ml ampicillin, 15 μ g/ml chloramphenicol, 10 μ g/ml kanamycin, 10 μ g/ml hemin and 50 μ g/ml aminolevulinic acid, cultured

under dark conditions or light conditions according to the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 11.

Table 11

		Relative absorbance			
E. coli strain	Culture conditions	Concentration of test			
		10 ppm	1 ppm	0 ppm	
BT3/pACYCSP+pTVHVF1	in the light	0.39	0.94	1.0	
BT3/pACYCSP+pTVHVF1	in the dark	0.94	0.96	1.0	
BT3/pACYCSP+pTV118N	in the light	0.12	0.24	1.0	
BT3/pACYCSP+pTV118N	in the dark	0.80	0.91	1.0	

Example 21

Introduction of the Gene Encoding Variant Barley

A plasmid for introducing the gene encoding barley ferrochelatase into tobacco by Agrobacterium infection method was constructed. The plasmid pTVHVF1 described in Example 20 was digested with restriction enzyme Nco I followed by blunting the DNA with DNA polymerase I by

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adding nucleotides to the double-stranded DNA gap. the DNA was dephosphorylated with alkaline phosphatase derived from calf intestine and cyclized by inserting phosphorylated BamHI linker (4610P, manufactured by Takara Shuzo Co., Ltd.) to construct plasmid pTVHVF2. pTVHVF2 was digested with restriction enzyme EcoRI followed by blunting of the DNA with DNA polymerase I by adding nucleotides to the double-stranded DNA gap. Further, the DNA was dephosphorylated with alkaline phosphatase derived from calf intestine and cyclized by inserting phosphorylated SalI linker (4680P, manufactured by Takara Shuzo Co., Ltd.) to construct plasmid pTVHVF3. Plasmid pBI121KS prepared in Example 9 was digested with restriction enzymes BamHI and SalI to remove B-glucuronidase gene. The DNA fragment gene encoding containing the the variant ferrochelatase was prepared by digesting the above pTVHVF3 with restriction enzymes BamHI and SalI. The resultant DNA fragment was inserted into plasmid pBI121KS with replacing β-glucuronidase gene to prepare plasmid pBIHVF (Fig. 13) in which variant barley gene joined downstream from 35S promoter.

The plasmid pBIHVF was introduced into Agrobacterium tumefaciens LBA4404. The resultant transformants were cultured in a medium containing 300 µg/ml streptomycin, 100 µg/ml rifampicin and 25 µg/ml

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kanamycin, followed by selecting the desired transformants to isolate *Agrobacterium* strain bearing pBIHVF.

Leaf pieces of tobacco cultured sterilely were infected with said Agrobacterium strain and, according to the same manner as in Example 5, tobacco into which the gene encoding the variant barley ferrochelatase was introduced was obtained.

Example 22

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Gene Encoding
Variant Barley Ferrochelatase

The level of resistance to the PPO inhibitory-type herbicidal compound represented by Structure 8 was confirmed quantitatively by testing tobacco into which the gene encoding the variant barley ferrochelatase prepared in Example 21 was introdued according to the same manner as in Example 6. The results obtained from 4 clones (HVF 1-4) of tobacco introduced with the gene encoding the variant barley ferrochelatase and control recombinant tobacco are shown in table 12. In the table, the resistant levels to the herbicidal compound are represented by percentages of the total chlorophyll content of leaf pieces treated with herbicidal compound to that of untreated leaf pieces.

Table 12

Recombinant		Total chlorophyll content (mg/ g-fresh weight)		
tobacco	untreated- leaf	treated-leaf	level to test compound(%)	
control	1.93	0.160	8.29	
HVF-1	0.876	0.930	106	
HVF-2	1.14	1.16	102	
HVF-3	1.06	1.04	98.1	
HVF-4	1.48	1.42	95.9	

Example 23

Test of Variant Protein of Cucumber Ferrochelatase Having Specific Affinity for Protoporphyrin IX for Capability of Giving Resistance to Herbicidal Compounds

using cucumber by PCR was carried out ferrochelatase cDNA clone isolated by the method described in Miyamoto, K. et al., Plant Physiol., 105; p 769 (1994) template, an oligonucleotide composed of nucleotide sequence of SEQ ID NO: 21 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 22 as primers to amplify the DNA fragment encoding cucumber sequence had been deleted ferrochelatase whose signal variant cucumber (hereinafter referred to as the ferrochelatase). The oligonucleotides were prepared with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA oligonucleotides synthesizer) and purified with an

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purification cartridge (PE Applied Biosystems; OPC cartridge). The PCR was carried out by repeating a cycle for maintaining at 94°C for 1 minute, at 55°C for 2 minutes and then at 72°C for 3 minutes 30 times. The amplified DNA fragments were digested with restriction enzymes BamHI and SacI, and inserted between BamHI restriction site and SacI restriction site of plasmid pTV119N (manufactured by Takara Shuzo Co., Ltd.) to construct plasmid pTVCSF (Fig. 14).

The plasmids pTVCSF and pTV118N were introduced into E. coli BT3/pACYCSP strain prepared in Example 3 respectively according to the method described in Hanahan, D.J., Mol. Biol., 166; p 557 (1983). E. coli BT3/pACYCSP+pTVCSF strain bearing plasmid pACYCSP and pTVCSF, and E. coli BT3/pACYCSP+pTV118N strain bearing plasmid pACYCSP and pTV118N were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin.

These E. coli strains were inoculated into YPT medium containing 10 or 1 ppm of the PPO inhibitory-type herbicidal compound represented by Structure 8, 100 µg/ml ampicillin, 15 µg/ml chloramphenicol, 10 µg/ml kanamycin, 10 µg/ml hemin and 50 µg/ml aminolevulinic acid, cultured under dark conditions or light conditions according to the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of liquid culture

medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 13.

Table 13

		Relative absorbance			
E. coli strain	Culture conditions	Concentration of test			
		10 ppm	1 ppm	0 ppm	
BT3/pACYCSP+pTVCSF	in the light	0.73	0.78	1.0	
BT3/pACYCSP+pTVCSF	in the dark	0.89	0.92	1.0	
BT3/pACYCSP+pTV118N	in the light	0.06	0.08	1.0	
BT3/pACYCSP+pTV118N	in the dark	0.81	0.91	1.0	

Example 24

Introduction of the Gene Encoding Variant Cucumber Ferrochelatase into Tobacco

A plasmid for introducing the gene encoding the modified cucumber ferrochelatase into tobacco by Agrobacterium infection method was constructed. Plasmid pBI121 (manufactured by Colntech) was digested with restriction enzymes BamHI and SacI to remove $\beta\text{--glucuronidase}$ gene. A DNA fragment containing the gene encoding the variant cucumber ferrochelatase was prepared by digesting plasmid pTVCSF described in Example 23 with restriction

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enzymes BamHI and SacI. The resultant DNA fragment was introduced into plasmid pBI121 with replacing β -glucuronidase gene to prepare plasmid pBICSF (Fig. 15) in which variant cucumber ferrochelatase gene was joined downstream from 35S promoter.

The plasmid pBICSF was introduced into Agrobacterium tumefaciens LBA4404. The resultant transformants were cultured in a medium containing 300 $\mu g/ml$ streptomycin, 100 $\mu g/ml$ rifampicin and 25 $\mu g/ml$ kanamycin, followed by selecting the desired transformants to isolate Agrobacterium strain bearing pBICSF.

Leaf pieces of tobacco cultured sterilely were infected with said Agrobacterium strain to obtain tobacco introduced with the gene encoding the modified cucumber ferrochelatase according to the same manner as in Example 5.

Example 25

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Gene Encoding
Variant Cucumber Ferrochelatase

The level of resistance to PPO inhibitory-type herbicidal compounds is confirmed quantitatively by testing tobacco introduced with the gene encoding the modified cucumber ferrochelatase prepared in Example 24 according to the same manner as in Example 6.

Example 26

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Isolation of E. coli Coproporphyrinogen III
Oxidase (hemF) Gene

Genomic DNA was prepared from E. coli LE392 strain using Kit ISOPLANT for genome DNA preparation (manufactured by Nippon Gene). An oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 23 and an oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 24 were synthesized according to nucleotide sequences of its 5' and 3' regions of E. coli hemF gene X754131. registered in (Accession GenBank oligonucleotides were prepared with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotides purification cartridge (PE Applied Biosystems; OPC cartridge). PCR was carried out by using about 1 µg of E. coli LE392 strain genomic DNA as a template and the above oligonucleotides (each 10 pmol) as primers to amplify the DNA fragment containing E. coli hemF gene. The PCR was carried out by repeating a cycle for maintaining at 96°C for 1 minute, at 55°C for 2 minutes and then at 72°C for 3 minutes 30 times.

Example 27

The DNA fragment containing hemF gene amplified by the method described in Example 26 was digested with

restriction enzymes FbaI and PstI, and inserted between BamHI restriction site and PstI restriction site of commercially available plasmid pUCll®N (manufactured by Takara Shuzo Co., Ltd.) to construct plasmid pHEMF (Fig. 16).

The plasmid pHEMF and pTV118N were introduced into E. coli BT3/pACYCSP strain prepared in Example 3 respectively according to the method described in Hanahan, D. J., Mol. Biol., 166; p 557 (1983). E. coli BT3/pACYCSP+pHEMF strain bearing plasmid pACYCSP and pHEMF, and E. coli BT3/pACYCSP+pTV118N strain bearing plasmid pACYCSP and pTV118N were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin.

These E. coli strains were inoculated into YPT medium containing 10 or 1 ppm of the PPO inhibitory-type herbicidal compound represented by Structure 8, 100 µg/ml ampicillin, 15 µg/ml chloramphenicol, 10 µg/ml kanamycin, 10 µg/ml hemin and 50 µg/ml aminolevulinic acid, cultured under dark conditions or light conditions according to the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are

shown in Table 14.

Table 14

		Relative absorbance Concentration of test compound			
E. coli strain	Culture conditions				
		10 ppm	1 ppm	0 ppm	
BT3/pACYCSP+pHEMF	in the light	0.48	1.0	1.0	
BT3/pACYCSP+pHEMF	in the dark	0.94	0.95	1.0	
BT3/pACYCSP+pTV118N	in the light	0.06	0.16	1.0	
BT3/pACYCSP+pTV118N	in the dark	0.96	0.98	1.0	

Example 28

Introduction of E. coli hemF gene into Tobacco

A plasmid for introducing E. coli hemF gene into a plant by Agrobacterium infection method was constructed. First, for obtaining E. coli hemF gene, an oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 25 and an oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 26 were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA oligonucleotide purified with an synthesizer) and Biosystems; OPC purification cartridge (PE Applied carried out by using the PCR was cartridge). oligonucleotide primers according to the same manner as in Example 26 to amplify the DNA fragment containing E. coli hemF gene.

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Plasmid pBI121 (manufactured by Clontech) was digested with restriction enzymes BamHI and SacI to remove β -glucuronidase gene. The DNA fragment containing the gene encoding the E. coli hemF gene was prepared by digesting DNA fragment with restriction the above PCR-amplified The resultant DNA fragment was enzymes BamHI and SacI. into plasmid pBI121 with replacing introduced glucuronidase gene to prepare plasmid pBIHEMF (Fig. 17) in which E. coli hemF gene was joined downstream from 35S promoter.

The plasmid pBIHEMF was introduced into Agrobacterium tumefaciers LBA4404. The resultant transformants were cultured in a medium containing 300 $\mu g/ml$ streptomycin, 100 $\mu g/ml$ rifampicin and 25 $\mu g/ml$ kanamycin, followed by selecting the desired transformants to isolate Agrobacterium strain bearing pBIHEMF.

Leaf pieces of tobacco cultured sterilely were infected with the Agrobacterium strain to obtain tobacco introduced with E. coli hemF gene according to the same manner as in Example 5.

Example 29

Compounds of Tobacco Introduced with the E. coli hemF Gene
The level of resistance to the PPO inhibitory-type
herbicidal compounds is confirmed quantitatively by testing

tobacco introduced with the *E. coli* hemF gene (prepared in Example 28) according to the same manner as in Example 6.

Example 30

Binding Test of Porphyrin Compound-Binding

5 Protein to Protoporphyrin IX

A phage library presenting a protein containing an amino acid sequence composed of 5 random amino acids and a phage clone displaying a protein containing an amino acid sequence HASYS or RASSL (wherein H is histidine, A is alanine, S is serine, Y is tyrosine, R is arginine and L is leucine) which can specifically bind to porphyrin compound 5, 10, 15, 20-tetrakis (N-methylpyridinium-4-y1)-21H, 23H-porphine (H₂TMpyP) were prepared according to the method described in KITANO et al., Nihon Kagakukai (Chemical Society of Japan) 74th Spring Annual Meeting Pre-Published Abstracts of Presentation II, p 1353, 4G511 (1998).

First, the phage library displaying a protein containing an amino acid sequence composed of 5 random amino acids was constructed. Mixed oligonucleotides composed of the nucleotide sequence of SEQ ID NO: 27 and mixed oligonucleotides composed of the nucleotide sequence of SEQ ID NO: 28 were synthesized. The mixed oligonucleotides were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotide purification cartridge (PE

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Applied Biosystems; OPC cartridge). The above mixed oligonucleotides (each 50 pmol) were phosphorylated at 5' end by treating with T4 DNA kinase respectively. They were mixed and, after heating at 70°C for 10 minutes, subjected to annealing by cooling slowly to room temperature at rate Plasmid pCANTAB5E 0.5°C/minute. (manufactured Pharmacia Biotech) was digested with restriction enzymes Sfil and Notl to remove the recombinant antibody gene ScFv. The above phosphorylated and annealed oligonucleotide pair was inserted into the portion of the above recombinant antibody gene ScFv to prepare a plasmid containing a nucleotide sequence encoding a protein composed of a 5 acid sequence upstream from random amino comprising an amino acid sequence of M13 phage coat protein. The plasmid was introduced into E. coli TG-1 strain according to the method described in Hanahan, D.J., Mol. Biol. 166; p 557 (1983) and cultured in 2 x YT medium (10 q/liter yeast extract, 15 g/liter tryptone and 5 g/liter NaCl, pH 7.2) containing 100 μ g/ml ampicillin to obtain recombinant E. coli TG-1 strain. The recombinant E. coli TG-1 strain was inoculated into 2 x YT medium containing 100 µg/ml ampicillin and cultured with shaking at 37°C. Then, 1 hour after initiation of culture, 6 x 10^{10} pfu helper-phage M13K07 (manufactured by Pharmacia Biotech) was inoculated to the medium, and culture was continued for additional 18 DOROTYTO . 102700

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hours with shaking. Then, the liquid culture medium was centrifuged at $1,000 \times g$ for 20 minutes to collect the phage library displaying a protein containing the amino acid sequence composed of 5 random amino acids.

For preparing the phage clone displaying a protein containing the amino acid sequence HASYS (SEQ ID NO: 53), an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 29 and an oligonucleotide composed the nucleotide sequence of SEQ ID NO: synthesized. And, for preparing the phage clone displaying a protein containing the amino acid sequence RASSL (SEQ ID No: 55), an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 31 and an oligonucleotide composed nucleotide sequence of SEQ ID NO: 32 These oligonucleotides were synthesized with synthesized. a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA purified with an oligonucleotide synthesizer) and purification cartridge (PE Applied Biosystems; OPC The phage clone displaying the protein cartridge). containing the amino acid sequence HASYS or RASSL was obtained by the same operation as the above that obtaining the phage library displaying a protein containing the amino acid sequence composed of 5 random amino acids.

A phage suspension containing the phage clone displaying the protein containing the amino acid sequence

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HASYS, the phage clone displaying the protein containing the amino acid sequence RASSL or the phage library displaying the protein containing the amino acid sequence consisting of 5 random amino acids (titer $10^5\ \text{pfu}$) was nitro cellulose filter respectively spotted to (manufactured by Schleicher & Schuell), and then the nitro cellulose filter was blocked by shaking it in PBT buffer (137 mM NaCl, 8.10 mM Na, HPO, 2.68 mM KCl, 1.47 mM KH, PO, 0.05% Tween 20, pH 7.2) containing 1% bovine serum albumin. The nitro cellulose filter was washed with PBT buffer and shaken for 18 hours in 2 x SSC buffer (0.3 M NaCl, 0.03M sodium citric acid) containing 10 µM protoporphyrin IX. Further, said nitro cellulose filter was washed with 2 x dried, and fluorescence derived buffer, protoporphyrin IX was detected under ultraviolet light (365 nm).

The spots of the phage library did not show fluorescence, while the spots of both phage clones displaying the protein containing the amino acid sequence HASYS and that containing the amino acid sequence RASSL showed clear fluorescence.

Example 31

Test of Protoporphyrin IX Binding Protein for Capability of Giving Resistance to Herbicidal Compounds

First, a plasmid which could express the gene

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encoding the protein containing the amino acid sequence HASYS (SEO ID NO: 53), or the amino acid sequence RASSL (SEQ ID NO: 55) was prepared. For preparing the plasmid expressing the gene encoding the protein capable of composed of the amino acid sequence of SEQ ID NO: (hereinafter referred to as the protein MGHASYS), oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 33 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 34 were synthesized. oligonucleotides were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotide purification cartridge (PE Biosystems; OPC cartridge). The above Applied oligonucleotides (each 50 pmol) were phosphorylated at 5' end by treating with T4 DNA kinase, respectively. were mixed and then, after heating for 10 minutes at 70°C, subjected to annealing by cooling slowly to temperature at rate of 0.5°C/minute. Plasmid pTV118N was digested with restriction enzymes NcoI and EcoRI to remove the gene fragment consisting of 16 base pairs. pHASYS capable of expressing the gene encoding protein MGHASYS was prepared by inserted the above phosphorylated and annealed oligonucleotide pairs into the position of the above 16 base pairs.

Then, for preparing the plasmid capable of

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expressing the gene encoding the protein consisting of amino acid sequence of SEQ ID NO: 56 (hereinafter referred to as protein MGRASSL), an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 35 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 36 were synthesized. The oligonucleotides were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA oligonucleotide purified with an synthesizer) and cartridge (PE Applied Biosystems; purification cartridge). Plasmid pRASSL capable of expressing the gene encoding protein MGRASSL was prepared by the same procedure as that for plasmid pHASYS.

A plasmid capable of expressing the gene encoding the protein containing the amino acid sequence YAGY or YAGF (wherein Y is tyrosine, A is alanine, G is glycine, F is phenylalanine) (Sugimoto, N., Nakano. S., Chem. Lett. p 939, 1997) capable of binding to porphyrin compound H2TMPyP was prepared. For preparing the plasmid capable of expressing the gene encoding the protein consisting of the amino acid sequence of SEQ ID NO: 58 (hereinafter referred to as an oligonucleotide composed protein MGYAGY), nucleotide sequence of SEQ ID NO: 37 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 38 were preparing the plasmid capable of synthesized. For expressing the gene encoding the protein composed of the amino acid sequence of SEQ ID NO: 60 (hereinafter referred to as protein MGYAGF), an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 39 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 40 were also synthesized. These oligonucleotides were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotide Applied Biosystems; purification cartridge (PE Plasmid pYAGY capable of expressing the gene cartridge). encoding the protein MGYAGY and plasmid pYAGF capable of expressing the gene encoding protein MGYAGF were prepared by the same procedure as that for plasmid pHASYS.

The above plasmids pHASYS, pRASSL, pYAGY, pYAGF and pTV118N were introduced into E. coli BT3/pACYCSP strain prepared in Example 3 respectively according to the method described in Hanahan, D. J., Mol. Biol., 166; p 557 (1983). E. coli BT3/pACYCSP+pHASYS strain bearing plasmid pACYCSP and pHASYS, E. coli BT3/pACYCSP+pRASSL strain bearing plasmid pACYCSP and pRASSL, E. coli BT/pACYCSP+pYAGY strain pYAGY, E . colibearing plasmid pACYCSP and BT3/pACYCSP+pYAGF strain bearing plasmid pACYCSP and pYAGF and E. coli BT3/pACYCSP+pTV118N strain bearing plasmid pACYCSP and pTV118N were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 ug/ml chloramphenicol and 10 µg/ml kanamycin.

These *E. coli* strains were inoculated into YPT medium containing 1 ppm of the PPO inhibitory-type herbicidal compound represented by Structure 8, 100 µg/ml ampicillin, 15 µg/ml chloramphenicol, 10 µg/ml kanamycin, 10 µg/ml hemin and 50 µg/ml aminolevulinic acid, cultured under dark conditions or light conditions according to the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of liquid culture medium was measured at 600 nm. By taking the absorbance of the medium without the herbicidal compound as 1, the relative value of the absorbance of the medium containing the herbicidal compound was calculated. The results are shown in Table 15.

Table 15

		Relative	absorbance
E. coli strain	Culture conditions		
		1 ppm	0 ppm
BT3/pACYCSP+pHASYS	in the light	0.65	1.0
BT3/pACYCSP+pHASYS	in the dark	0.96	1.0
BT3/pACYCSP+pRASSL	in the light	0.59	1.0
BT3/pACYCSP+pRASSL	in the dark	1.0	1.0
BT3/pACYCSP+pYAGY	in the light	0.48	1.0
BT3/pACYCSP+pYAGY	in the dark	0.99	1.0
BT3/pACYCSP+pYAGF	in the light	0.62	1.0
BT3/pACYCSP+pYAGF	in the dark	0.96	1.0

втз	/pACYCSP+pTV118N	in	the	light	0.07	1.0
втз	/pACYCSP+pTV118N	in	the	dark	0.93	1.0

Further, a plasmid capable of expressing a gene encoding a protein containing an amino acid sequence in which one unit of the amino acid sequences HASYS or RASSL were repeatedly joined. For preparing the plasmid capable of expressing the gene encoding the protein composed of the amino acid sequence of SEO ID NO: 61 (hereinafter referred as protein MG(HASYS), (HASYS), referred to as sequence in which peptide HASYS was repeatedly joined to each other n times), an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 41, SEQ ID NO: 42, SEQ ID 43 or SEQ ID NO: 44 was synthesized. These NO: oligonucleotides were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotide purification cartridge (PE Biosystems; OPC cartridge). First, oligonucleotide composed of the nucleotide sequence of SEQ TD NO. and the oligonucleotide composed of nucleotide sequence of SEQ ID NO: 43 were phosphorylated respectively at 5' end by treating with T4 DNA kinase. Thereafter, the oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 41 and the oligonucleotide composed of the phosphorylated nucleotide sequence of SEQ ID NO: 42

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or the oligonucleotide composed of the phosphorylated sequence of SEQ ID NO: 43 nucleotide oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 44 were mixed (each 300 pmol), and after heating for 5 minutes at 70°C, annealed by cooling slowly to room temperature at rate of 0.5°C/minute. The above two annealed oligonucleotide pairs were mixed and ligated with T4 DNA ligase, then the resultant DNA fragment was phosphorylated with T4 DNA kinase at 5' end. On the other hand, vector pTV118N was digested with restriction enzymes NcoI and EcoRI to remove a DNA fragment of 16 base pairs and the above phosphorylated DNA fragment was inserted into this portion to obtain plasmid pHASYS4 expressing the gene encoding protein MG(HASYS) 4.

Further, for preparing the plasmid capable of expressing the gene encoding the protein composed of the amino acid sequence of SEQ ID NO: 62 (hereinafter referred to as protein MG(HASYS), an oligonucleotide composed of the nucleotide sequence of SEO ID NO: 45 and oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 46 were synthesized. These oligonucleotides were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotide purification cartridge (PE Applied cartridge). First, the above Biosystems; OPC

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oligonucleotides were phosphorylated at 5' end by treating T4 DNA kinase. Thereafter, an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 41 and an oligonucleotide composed of the phosphorylated nucleotide sequence of SEO ID NO: 42 were mixed (each 300 pmol), an oligonucleotide composed of the phosphorylated nucleotide sequence of SEQ ID NO: 43 and an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 44 were mixed (each 300 pmol), and further, an oligonucleotide composed of the phosphorylated nucleotide sequence of SEQ ID NO: 45 and an oligonucleotide composed of the phosphorylated nucleotide sequence of SEQ ID NO: 46 were mixed (each 600 pmol). These three mixtures were heated for 5 minutes at 70°C, and annealed by cooling slowly to room temperature at rate of 0.5°C/minute, respectively. The above annealed oligonucleotide pairs were mixed, and ligated with T4 DNA ligase, and then the resultant DNA fragment was phosphorylated with T4 DNA kinase at 5' end. capable of expressing protein MG(HASYS)8 pHASYS8 prepared in the same manner as that for the above plasmid pHASYS4.

Then, for preparing a plasmid capable of expressing the gene encoding the protein composed of the amino acid sequence of SEQ ID NO: 63 (hereinafter referred to as protein MG(RASSL),, (RASSL), referred to as a

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sequence in which peptide RASSL was repeatedly joined to each other n times), an oligonucleotide composed of the nucleotide sequence of SEO ID NO: 47, SEQ ID NO: 48, SEQ ID NO: 49 or SEQ ID NO: 50 were synthesized. Also, for preparing a plasmid capable of expressing the gene encoding the protein composed of the amino acid sequence of SEQ ID NO: 64 (hereinafter referred to as protein MG(RASSL) 8), an oligonucleotide composed of the nucleotide sequence of SEQ ID NO: 51 and an oligonucleotide composed of the nucleotide sequence of SEQ ID No: 52 were synthesized. These oligonucleotides were synthesized with a DNA synthesizer (PE Applied Biosystems; Model 394 DNA/RNA synthesizer) and purified with an oligonucleotide purification cartridge (PE Applied Biosystems; OPC cartridge).

Plasmid pRASSL4 capable of expressing protein MG(RASSL), were prepared according to the same manner as that for the above plasmid pHASYS4. Plasmid pRASSL8 capable of expressing protein MG(RASSL), were also prepared according to the same manner as that for the above plasmid pHASYS8.

The above plasmids pHASYS4, pHASYS8, pRASSL4, pRASSL4 and pTV118N were introduced into E. coli BT3/pACYCSP strain prepared in Example 3 respectively according to the method described in Hanahan, D. J., Mol. Biol., 166; p 557 (1983). E. coli BT3/pACYCSP+pHASYS4

strain bearing plasmid pACYCSP and pHASYS4, *E. coli* BT3/pACYCSP+pHASYS8 strain bearing plasmid pACYCSP and pHASYS8, *E. coli* BT3/pACYCSP+pRASSL4 strain bearing plasmid pACYCSP and pRASSL4, *E. coli* BT3/pACYCSP+pRASSL8 strain bearing plasmid pACYCSP and pRASSL8 and *E. coli* BT3/pACYCSP+pTV118N strain bearing plasmid pACYCSP and pTV118N were obtained by culturing the above strains in YPT medium containing 100 µg/ml ampicillin, 15 µg/ml chloramphenicol and 10 µg/ml kanamycin.

These *E. coli* strains were inoculated into YPT medium containing 1 ppm of the PPO inhibitory-type herbicidal compound represented by Structure 8, 100 µg/ml ampicillin, 15 µg/ml chloramphenicol, 10 µg/ml kanamycin, 10 µg/ml hemin and 50 µg/ml aminolevulinic acid, cultured under dark conditions or light conditions according to the same manner as in Example 2. Then, 18 hours after initiation of culture, the absorbance of the liquid culture medium was measured at 600 nm. By taking the absorbance of the culture medium without the herbicidal compound as 1, the relative value of the absorbance of the culture medium containing the herbicidal compound was calculated. The results are shown in Table 16.

Table 16

E. coli strain		Relative absorbance	
	Culture condition	Concentration of test compound	
		1 ppm	0 ppm
BT3/pACYCSP+pHASYS4	in the light	0.91	1.0
BT3/pACYCSP+pHASYS4	in the dark	1.0	1.0
BT3/pACYCSP+pHASYS8	in the light	0.57	1.0
BT3/pACYCSP+pHASYS8	in the dark	1.0	1.0
BT3/pACYCSP+pRASSL4	in the light	1.1	1.0
BT3/pACYCSP+pRASSL4	in the dark	0.98	1.0
BT3/pACYCSP+pRASSL8	in the light	0.79	1.0
BT3/pACYCSP+pRASSL8	in the dark	1.0	1.0
BT3/pACYCSP+pTV118N	in the light	0.15	1.0
BT3/pACYCSP+pTV118N	in the dark	0.81	1.0

Example 32

 $\hbox{ Introduction of the Gene Encoding Protoporphyrin } \\ \hbox{IX Binding Peptide into Tobacco} \\$

A plasmid for introducing the gene encoding the protoporphyrin IX binding peptide into tobacco by Agrobacterium method was constructed. The plasmid pHASYS8 prepared in Example 31 was digested with restriction enzyme NcoI followed by blunting the DNA with DNA polymerase I with addition of nucleotides to the double-stranded DNA gap. Then, the DNA was dephosphorylated with alkaline phosphatase derived from calf intestine and cyclized by inserting phosphorylated BamH I linker (4610P, manufactured by Takara Syuzo Co., Ltd.) to construct plasmid pHASYS8B. Plasmid

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(manufactured by Clonetech) was digested with restriction enzymes BamHI and SacI to remove β-glucuronidase gene. On the other hand, plasmid pHASYS8B was digested with restriction enzymes BamHI and SacI to prepare the DNA fragment containing the gene encoding protein MG(HASYS) a, the resultant DNA fragment was inserted into plasmid pBI121 with replacing β-glucuronidase gene to prepare plasmid which the gene pBIHASYS8 (Fig. 18) in protoporphyrin IX binding protein MG(HASYS), was joined downstream from 35S promoter.

A plasmid for introducing the gene encoding the protoporphyrin IX binding peptide MG(RASSL)₈ into a plant by Agrobacterium infection method was constructed. Plasmid pBIRASSL8 (Fig. 19) in which the gene encoding protoporphyrin IX binding protein MG(RASSL)₈ was joined downstream from 35S promoter was prepared from pRASSL8 according to the same procedure as that for pBIHASYS8.

The above plasmid pBIHASYS8 and pBIRASSL8 were introduced into Agrobacterium tumefaciens LBA4404 respectively. The resultant transformants were cultured in a medium containing 300 µg/ml streptomycin, 100 µg/ml rifampicin and 25 µg/ml kanamycin, followed by selecting the desired transformants to isolate Agrobacterium strains bearing pBIHASYS8 and pBIRASSL8, respectively.

Leaf pieces of tobacco cultured sterilely are

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infected with said Agrobacterium strains to obtain tobacco introduced with the gene encoding protoporphyrin IX binding protein MG(HASYS), and the tobacco introduced with the gene encoding protoporphyrin IX binding protein MG(RASSL), in the same manner as in Example 5.

Example 33

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Gene Encoding the
Protoporphyrin IX Binding Peptide

The level of resistance to herbicidal compounds is confirmed quantitatively by testing tobacco introduced with the gene encoding the protoporphyrin IX binding peptide prepared in Example 32 according to the same manner as in Example 14.

Example 34

Isolation of PPO Gene of Arabidopsis thaliana Having Herbicidal Compound-Resistant Mutation

A plasmid containing PPO gene of Arabidopsis thaliana obtained by the method described by Narita, S. et al., Gene, 182; p 169 (1996) was digested with the restriction enzyme NcoI, and nucleotides were added to the gap of the double-stranded DNA by using DNA Polymerase I to blunt the end of the DNA. Then, the 5'-end of the DNA was dephosphorylated with an alkaline phosphatase derived from calf small intestine, followed by insertion of a

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phosphorylated BamHI linker (4810P manufactured by Takara Shuzo Co., Ltd.) therein and cyclization to construct plasmid pAGE17B. The plasmid pAGE17B was digested with BamHI and SacI to obtain a gene fragment containing PPO gene of Arabidopsis thaliana. The fragment was inserted between BamHI and SacI of a commercially available vector, pKF19k for site-directed mutagenesis (manufactured by Takara Shuzo Co., Ltd.), to construct plasmid pKFATP.

Then, for conversion of the 220th alanine into valine, a PPO inhibitory-type herbicidal compound-resistant mutation in PPO protein of Arabidopsis thaliana disclosed in WO 9534659, base substitution (substitution of "T" for the 659 base "C") of DNA was introduced into the above PPO gene of Arabidopsis thaliana. First, an oligonucleotide primer for mutagenesis represented by SEQ ID NO: 65 was The oligonucleotide primer was synthesized synthesized. with a DNA synthesizer (PE Applied Biosystems: Model 394 DNA/RNA Synthesizer) and purified with an oligonucleotide purification cartridge (PE Applied Biosystems: OPC Cartridge). The 5'-end of the oligonucleotide primer was phosphorylated with T4 DNA kinase. According to the manual attached to a commercially available site-directed mutagenesis kit, Mutan-Super Express Km (manufactured by Takara Shuzo Co., Ltd.), a reaction mixture containing 10 ng of the above plasmid pKFATP as template DNA, 5 pmol of

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the attached selection primer, 5 pmol of the above phosphorylated oligonucleotide primer for mutagenesis and the like was prepared and PCR was carried out. The PCR was carried by repeating a cycle for maintaining at 94°C for 1 minute, at 55°C for 1 minute and then at 72°C for 3 minutes 30 times. The resultant reaction mixture was purified by ethanol precipitation and the precipitate was dissolved in 5 ul of sterilized-distilled water. According to the attached manual, its 2 µl portion was used for introduction into a commercially available E. coli competent cell, MV1184 (manufactured by Takara Shuzo Co., Ltd.), and plated on LB agar culture medium (1% tryptone, 0.5% yeast extract, 0.5% NaCl, 1% agar) containing 50 µg/ml of kanamycin. After incubation at 37°C, the resultant clone was cultured in LB liquid medium containing 50 µg/ml of kanamycin to prepare plasmid DNA. The introduction of the desired herbicidal compound-resistant mutation A220V was confirmed by analyzing the nucleotide sequence of the PPO gene of Arabidopsis thaliana having herbicidal compound-resistant mutation contained in the resultant plasmid pKFATP1.

Example 35

Introduction of PPO Gene of Arabidopsis thaliana
Having Herbicidal Compound-Resistant Mutation into Tobacco

A plasmid was constructed for introducing the PPO gene of Arabidopsis thaliana having herbicidal compound-

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resistant mutation (hereinafter referred to as Arabidopsis thaliana PPO(A220V) gene) into a plant by Agrobacterium infection method. Binary vector pBI121 (manufactured by Clontech) was digested with the restriction enzymes BamHI and SacI to remove β -glucuronidase gene. On the other hand, the plasmid pKFATP1 described in Example 34 was digested with restriction enzymes BamHI and SacI to prepare a DNA fragment containing Arabidopsis thaliana PPO(A220V) gene. Instead of the above β -glucuronidase gene, the resultant DNA fragment was inserted in the binary vector pBI121 to construct the plasmid pNATP (Fig. 20).

The plasmid pNATP was introduced into Agrobacterium tumefaciens LBA4404 and this was cultured in LB culture medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pNATP.

Tobacco leaf pieces cultured sterilely were infected with the Agrobacterium strain and, according to the same manner as described in Example 5, tobacco bearing the introduced Arabidopsis thaliana PPO(A220V) gene was obtained.

Example 36

Production of Recombinant Tobacco Having

Arabidopsis thaliana PPO(A220V) Gene and Gene Encoding

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Variant Tobacco Chelatase Subunit

A plasmid was constructed for introducing both Arabidopsis thaliana PPO(A220V) gene and gene encoding a variant tobacco chelatase subunit into a plant Agrobacterium infection method. First, oligonucleotide primers composed of the nucleotide sequence represented by SEO ID NO: 66 and the nucleotide sequence represented by SEQ ID NO: 67, respectively, were synthesized. primers were synthesized bv DNA oligonucleotide synthesizer (PE Applied Biosystems: Model 394 purified with an oligonucleotide Synthesizer) and purification cartridge (PE Applied Biosystems: OPC Cartridge). PCR was carried out by using the primers and the plasmid pKFATP1 constructed in Example 34 as template DNA to prepare a DNA fragment containing Arabidopsis thaliana PPO(A220V) gene. The PCR was carried out by repeating a cycle for maintaining at 94°C for 1 minute, 55°C for 2 minutes and then 72°C for 3 minutes 30 times. The amplified DNA fragment was digested by the restriction HindIII and SalI to obtain a DNA fragment containing Arabidopsis thaliana PPO(A220V) gene. fragment was inserted between HindIII and SalI restriction sites of a commercially available vector, pUC19, construct plasmid pAP.

On the other hand, the plasmid pNG01 (Fig. 29)

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described in Shiota, N. et al., Plant Physiol., 106; p 17 (1994) was digested with the restriction enzyme HindIII and nucleotides were added to the gap of the double-stranded DNA with DNA Polymerase I to blunt the end of the DNA, followed by self-cyclization to construct the plasmid pNG04 The plasmid pNG04 was digested with the restriction enzyme XbaI to isolate a DNA fragment of about 1.1 kb composed of the terminator of a nopaline synthase The DNA fragment was inserted into the and 35S promoter. XhaI restriction site of the above plasmid pAP. selecting plasmid pAPNS wherein the terminator of the downstream nopaline synthase was ligated to Arabidopsis thaliana PPO(A220V) gene, digestion with the restriction enzymes HindIII and PstI was carried out to select a clone producing a DNA fragment of about 2.0 kb composed of Arabidopsis thaliana PPO(A220V) gene and the terminator of the nopaline synthase. Further, the plasmid pAPNS was digested with the restriction enzyme HindIII and nucleotides were added to the gap of the double-stranded DNA with DNA polymerase I to blunt the end of the DNA. 5'-end of the DNA was dephosphorylated with an alkaline phosphatase derived from calf small intestine phosphorylated KpnI linker (4668P manufactured by Takara was inserted therein, followed by Shuzo Co., Ltd.) cyclization to construct plasmid pAPNSK.

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plasmid pAPNSK was digested with The restriction enzymes KpnI and DraI to isolate a DNA fragment of about 2.8 kb composed of Arabidopsis thaliana PPO(A220V) gene, the terminator of the nopaline synthase located at the downstream of the gene and 35S promoter located at the downstream of the terminator. The fragment was inserted in KpnI restriction site of the plasmid pBITCHLH constructed in Example 9. For selecting the plasmid pBIAPTCH (Fig. 21) wherein Arabidopsis thaliana PPO(A220V) gene and a gene encoding a variant tobacco chelatase subunit were ligated to the downstream of 35S promoter, respectively, the resultant plasmid was digested with the restriction enzyme BamHI to select a clone producing a DNA fragment of about 2.8 kb composed of Arabidopsis thaliana PPO(A220V) gene, the terminator of the nopaline synthase and the promoter.

The plasmid pBIAPTCH was introduced into Agrobacterium tumefaciens LBA4404 and this was cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml rifampicin and 25 μ g/ml kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBIAPTCH.

Tobacco leaf pieces cultured sterilely were infected with the Agrobacterium strain and, according to the same manner as in Example 5, tobacco bearing the

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introduced both Arabidopsis thaliana PPO(A220V) gene and gene encoding the variant tobacco chelatase subunit was obtained.

Example 37

Confirmation of Resistance to Herbicidal Compound of Tobacco Bearing Introduced Arabidopsis thaliana PPO(A220V) Gene and Gene Encoding Variant Tobacco Chelatase Subunit

Leaves of the tobacco bearing the introduced both Arabidopsis thaliana PPO(A220V) gene and gene encoding the variant tobacco chelatase subunit produced in Example 36, those bearing the introduced Arabidopsis thaliana PPO(A220V) gene produced in Example 35, and the control recombinant tobacco leaves produced in Example 5 were collected, and each leaf was divided into the right and left equivalent pieces along the main vein. One of the pieces was treated with an aqueous solution containing 2.0 ppm PPO inhibitory-type herbicidal compound of Structure 8, while the other piece was not treated with the compound. These leaf pieces were placed on MS medium containing 0.8% agar and allowed to stand at room temperature for 7 days in a light place. Then, each leaf piece was ground in 5 ml of 80% aqueous acetone solution in a mortar with a pestle to extract chlorophyll. The extract was diluted 10 times with aqueous acetone solution and the absorbance was

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measured at 750 nm, 663 nm and 645 nm to calculate the total chlorophyll content according to the method described by Macknney G., J. Biol. Chem. (1941) 140, p 315. The resistant level to the herbicidal compound tested was represented by the percentage of the total chlorophyll content of the leave pieces treated with the herbicidal compound to that of untreated leaf pieces. The resistant level of the control recombinant tobacco was 2.88% and that of the tobacco bearing the introduced Arabidopsis thaliana PPO(A220V) gene was 12.2%. On the other hand, the resistant level of the tobacco bearing introduced both Arabidopsis thaliana PPO(A220V) gene and gene encoding the variant tobacco chelatase subunit was 61.6%.

Example 38

 ${\tt Isolation \ of \ Gene \ Encoding \ Chloroplast-Localized}$ ${\tt Type \ Ferrochelatase \ of \ } Arabidops is \ thaliana$

Total RNAs were prepared from leaf tissues of Arabidopsis thaliana ecotype WS by using RNeasy Plant Kit (manufactured by QIAGEN) according to the manual attached thereto. A DNA fragment containing a chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana was obtained by using RNA LA PCR Kit (AMV) Ver 1.1 (manufactured by Takara Shuzo Co., Ltd.) according to the manual attached thereto. First, 1st strand cDNA was synthesized by using Arabidopsis thaliana total RNAs as a

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template and Oligo dT-Adaptor Primer contained in the above kit as the primer with the reverse transcriptase contained in the above kit. Then, PCR was carried out by using the 1st strand cDNA as a template and LA Tag polymerase contained in the above kit to amplify a DNA fragment containing the gene encoding the chloroplast-localized type In this PCR, the ferrochelatase of Arabidopsis thaliana. primers used were the oligonucleotide primer composed of the nucleotide sequence of SEQ ID NO: 68 and the oligonucleotide primer composed of the nucleotide sequence of SEO ID NO: 69. The oligonucleotides were synthesized by a DNA synthesizer (PE Applied Biosystems: Model 394 DNA/RNA purified with oligonucleotide Synthesizer) and an Applied Biosystems: purification cartridge (PE Cartridge). The PCR was carried by maintaining at 94°C for 2 minutes and then repeating a cycle for maintaining at 94°C for 30 seconds, at 50°C for 30 seconds and then at 72°C for 7 minutes 30 times. After the PCR, the DNA fragment amplified by the PCR was cloned into the plasmid pCR2.1 by using TA Cloning Kit (manufactured by Invitrogen) according to the manual attached thereto. The resultant plasmid was digested with the restriction enzyme BamHI and analyzed by agarose gel electrophoresis. The plasmid from which 5.3 kb DNA fragment was detected was named pCRATF The plasmid has such a structure that the (Fig. 22).

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chloroplast-localized tvpe ferrochelatase gene Arabidopsis thaliana is inserted in DNA complementary to the lac promoter. When the nucleotide sequence of the chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana was analyzed, it agreed with the nucleotide sequence of the chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana described by Smith, A.G. et al., J. Biol. chem., 269; p 13405 (1994).

Example 39

Introduction of Gene Encoding Chloroplast-localized Type Ferrochelatase of Arabidopsis thaliana

A plasmid was constructed for introducing a gene chloroplast-localized type ferrochelatase encoding thaliana into plant by Agrobacterium Arabidopsis a infection method. First, the plasmid pCRATF constructed in Example 38 was digested with the restriction enzymes BamHI and SacI to prepare a DNA fragment containing chloroplastlocalized type ferrochelatase gene of Arabidopsis thaliana. Binary vector pBI121 (manufactured by Clontech) digested with the restriction enzymes BamHI and SacI to remove β -glucuronidase gene and, instead of this gene, the above DNA fragment containing the chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana inserted therein to construct the plasmid pBIATF (Fig. 23) wherein the ferrochelatase gene was ligated

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downstream of the 35S promoter.

The plasmid pBIATF was inserted into Agrobacterium tumefaciens LBA4404 and this was cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBIATF.

Tobacco leaf pieces cultivated sterilely are infected with the Agrobacterium strain and tobacco bearing the introduced chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana is obtained according to the same manner as in Example 5.

Example 40

Production of Recombinant Tobacco Bearing Both

Arabidopsis thaliana PPO(A220V) Gene and ChloroplastLocalized Type Ferrochelatase Gene of Arabidopsis thaliana

A plasmid is constructed for introducing both Arabidopsis thaliana PPO(A220V) gene and chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana into a plant by Agrobacterium infection method. First, the plasmid pAPNS constructed in Example 36 are digested with the restriction enzyme HindIII and nucleotides are added to the gap of the double-stranded DNA with DNA Polymerase I to blunt the end of the DNA. The 5'-end of the DNA is dephosphorylated with an alkaline phosphatase derived from

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calf small intestine and a phosphorylated BamHI linker (4610P manufactured by Takara Shuzo Co., Ltd.) is inserted therein, followed by cyclization to construct plasmid pAPNSB.

plasmid pAPNSB is digested with the The restriction enzymes BamHI and DraI to isolate a DNA fragment of about 2.8 kb composed of Arabidopsis thaliana PPO(A220V) gene, the terminator of nopaline synthase and The fragment is inserted in the BamHI 35S promoter. restriction site of the plasmid pBIATF constructed in Example 39. For selecting the plasmid pBIAPATF (Fig. 24) wherein Arabidopsis thaliana PP(A220V) gene and type ferrochelatase chloroplast-localized gene ofArabidopsis thaliana are ligated to the downstream of 35S promoter, the resultant plasmid is digested with restriction enzymes NotI and SacI to select a clone producing a DNA fragment of about 2.2 kb composed of the 35S promoter and the chloroplast-localized ferrochelatase gene of Arabidopsis thaliana.

The plasmid pBIAPATF is introduced into Agrobacterium tumefaciens LBA44044 and this is cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBIAPATF.

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Tobacco leaf pieces cultivated sterilely are infected with the Agrobacterium strain and tobacco bearing the introduced Arabidopsis thaliana PPO(A220V) gene and chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana is obtained according to the same manner as in Example 5.

Example 41

Comfirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Arabidopsis
thaliana PPO(A220V) Gene and Chloroplast-Localized Type
Ferrochelatase Gene of Arabidopsis thaliana

The levels of resistance to herbicidal compounds are confirmed quantitatively by testing the tobacco bearing the introduced Arabidopsis thaliana PPO(A220V) gene and chloroplast-localized type ferrochelatase gene of Arabidopsis thaliana produced in Example 40 according to the same manner as in Example 37.

Example 42

Isolation of Soybean Coproporphyrinogen III
Oxidase Gene

Total RNAs were prepared from leaf tissues of soybean (Glycine max cv. Jack) by using RNeasy Plant Kit (manufactured by QIAGEN) according to the manual attached thereto. Further, a DNA fragment containing a gene encoding soybean coproporphyrinogen III oxidase

2.0

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(hereinafter referred to as the present soybean CPOX) was obtained by using RNA LA PCR Kit (AMV) Ver 1.1 (manufactured by Takara Shuzo Co., Ltd.) according to the manual attached thereto. First, 1st strand cDNA was synthesized by using the soybean total RNAs as a template and Oligo dT-Adaptor Primer contained in the above kit as a primer with the reverse transcriptase contained in the above kit. Then, PCR was carried out by using the 1st strand cDNA as a template and LA Tag polymerase contained in the above kit to amplify a DNA fragment containing the present soybean CPOX gene. In this PCR, an oligonucleotide primer composed of the nucleotide sequence represented by SEQ ID NO: 70 and an oligonucleotide primer composed of the nucleotide sequence represented by SEQ ID NO: 71 were used. These oligonucleotides were synthesized by using a DNA synthesizer (PE Applied Biosystems: Model 394 DNA/RNA purified with an oligonucleotide Synthesizer) and purification cartridge (PE Applied Biosystems: OPC Cartridge). The PCR was carried out by maintaining at 94°C for 2 minutes and then repeating a cycle for maintaining at 94°C for 30 seconds, at 50°C for 30 seconds and then at 72°C for 7 minutes 30 times. After the PCR, the DNA fragment amplified by the PCR was cloned into plasmid pCR2.1 by using TA Cloning Kit (manufactured by Invitrogen) according to the manual attached thereto. The resultant

1.0

plasmid was digested with the restriction enzyme BamHI and analyzed by agarose gel electrophoresis. The plasmid from which 1.2 kb DNA fragment was detected was named pCRSCPOX (Fig. 25). The plasmid pCRSCPOX has such a structure that the present soybean CPOX gene is inserted into a DNA strand complementary to the lac promoter. When the nucleotide sequence of the DNA fragment in the plasmid was analyzed, it was confirmed to be the present soybean CPOX gene.

Example 43

Introduction of Present Soybean CPOX Gene into

A plasmid was constructed for introducing the present soybean CPOX gene into a plant by Agrobacterium infection method. First, the plasmid pCRSCPOX constructed in Example 42 was digested with the restriction enzymes BamHI and SalI to prepare a DNA fragment containing the present soybean CPOX gene. The plasmid pBI121KS constructed in Example 9 was digested with the restriction enzymes BamHI and SalI to remove β -glucuronidase gene and, instead thereof, the above DNA fragment containing the present soybean CPOX gene was inserted therein to construct the plasmid pBISCPOX (Fig. 26) wherein the gene was ligated to the downstream of 35S promoter.

The plasmid pBISCPOX was introduced into 25 Agrobacterium tumefaciens LBA4404 and it was cultured in LB

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medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an *Agrobacterium* strain bearing pBISCPOX.

Tobacco leaf pieces cultured sterilely is infected with the Agrobacterium strain and, according to the same manner as in Example 5, tobacco bearing the introduced present soybean CPOX gene is obtained.

Example 44

Production of Recombinant Tobacco Bearing

Arabidopsis thaliana PPO(A220V) Gene and Present Soybean

CPOX Gene

A plasmid is constructed for introducing both Arabidopsis thaliana PPO(A220V) gene and present soybean CPOX gene into a plant by Agrobacterium infection method. The plasmid pAPNSB constructed in Example 40 is digested with the restriction enzymes BamHI and DraI to isolate a DNA fragment of about 2.8 kb composed of Arabidopsis thaliana PPO(A220V) gene, the terminator of a nopaline synthase and 35S promoter. The fragment is inserted into the BamHI restriction site of the plasmid pBISCPOX constructed in Example 43. For selecting the plasmid pBIAPSCP (Fig. 27) wherein Arabidopsis thaliana PPO(A220V) gene and the present soybean CPOX gene are ligated to the downstream of the 35S promoter, respectively, the resultant

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plasmid is digested with the restriction enzymes NotI and SalI to select a clone producing a DNA fragment of about 2.0 kb composed of the 35S promoter and the present CPOX gene.

The plasmid pBIAPSCP is introduced into Agrobacterium tumefaciens LBA4404 and this is cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBIAPSCP.

Tobacco leaf pieces cultivated sterilely is infected with the Agrobacterium strain and, according to the same manner as in Example 5, tobacco bearing the introduced both Arabidopsis thaliana PPO(A220V) gene and present soybean CPOX gene is obtained.

Example 45

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Both Arabidopsis
thaliana PPO(A220V) Gene and Present Soybean CPOX Gene

The levels of resistance to herbicidal compounds are confirmed quantitatively by testing the tobacco bearing the introduced both Arabidopsis thaliana PPO(A220V) gene and present soybean CPOX gene produced in Example 44 according to the same manner as in Example 37.

Example 46

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Isolation of Glyphosate Resistant Gene

Glyphosate resistant sovbean (Glycine max) was seeded and cultivated at 27°C for 30 days. The first leaves of germinated individuals were collected, were frozen in liquid Nitrogen and were grounded in a mortar with a pestle. Genomic DNA was extracted from the ground material with a genomic DNA extracting reagent ISOPLANT (manufactured by NIPPON GENE) according to the manual attached thereto. PCR was carried out by using the genomic DNA as a template, an oligonucleotide primer composed of the nucleotide sequence represented by SEO ID NO: 72 and an oligonucleotide primer composed of the nucleotide sequence represented by SEQ ID NO: 73 to amplify a DNA fragment (hereinafter referred to as the present CTP-CP4 EPSPS gene) containing a nucleotide sequence encoding a chloroplast transit peptide sequence of EPSPS of petunia (Petunia hybrida) (hereinafter referred to as CTP) and EPSPS gene of Strain CP4). Agrobacterium (Agrobacterium sp. The oligonucleotides were synthesized by using DNA synthesizer (PE Applied Biosystems: Model 394 DNA/RNA purified with oligonucleotide Synthesizer) and an purification cartridge (PE Applied Systems: OPS Cartridge). The PCR was carried out by maintaining at 94°C for 5 minutes, 55°C for 2 minutes and then at 72°C for 3 minutes, and further repeating a cycle for maintaining at 94°C for 1

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minute, at 55°C for 2 minutes and then 72°C for 3 minutes 38 times, and, finally, further maintaining at 94°C for 1 minute, at 55°C for 2 minutes and then 72°C for 10 minutes. The amplified DNA fraction was ligated to a PCR product cloning site of plasmid pCR2.1 (manufactured Invitrogen), to construct the plasmid pCREPSPS (Fig. 28). Then, the plasmid was introduced in a competent cell of E. coli JM109 strain (manufactured by Takara Shuzo Co., Ltd.) to select an ampicillin resistant strain. The nucleotide sequence of the plasmid contained in the selected ampicillin resistant strain was determined by using Thermo Sequence II Dye Terminator kit (manufacture by Amersham Pharmacia Biotech) and DNA Sequencer 373S (manufactured by PE Biosystems). As a result, the nucleotide represented by SEO ID NO: 74 was revealed and it was confirmed that the plasmid pCREPSPS contained the present CTP-CP4 EPSPS gene.

Example 47

 $\qquad \qquad \text{Introduction of Present CTP-CP4} \ \ \text{EPSPS Gene into}$ $\ \ \text{Tobacco}$

A plasmid was constructed for introducing the present CTP-CP4 EPSPS gene into a plant by Agrobacterium infection method. First, pNG01 [Shiota et al., (1994) Plant Physiol., 106:17-23] (Fig. 29) was digested with the restriction enzyme HindIII and nucleotides were added to

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the gap of the double-stranded DNA with DNA Polymerase I to blunt the end of the DNA, followed by self-cyclization with T4 DNA ligase to obtain pNG04 (Fig. 30). The plasmid pNG04 was digested with the restriction enzyme XbaI to obtain a DNA fragment containing the terminator of a nopaline synthase and 35S promoter located at the downstream thereof. The fragment was inserted in the XbaI restriction site of plasmid pUC19 (manufactured by Takara Shuzo Co., Ltd.) to obtain pNT35S (Fig. 31). Then, the plasmid pCREPSPS constructed in Example 46 was digested with the restriction enzymes HindIII and SalI and the resultant DNA fragment containing the present CTP-CP4 EPSPS gene was inserted between HindIII and SalI restriction sites of pNT35S to obtain the plasmid pCENS (Fig. 32). The plasmid pCENS was digested with the restriction enzyme HindIII nucleotides were added to the gap of the double-stranded DNA with DNA Polymerase I to blunt the end of the DNA. 5'-end of the DNA was dephosphorylated with treatment of alkaline phosphatase derived from calf small intestine, followed by insertion of phosphorylated KpnI linker (4668A manufactured by Takara Shuzo Co., Ltd.) therein cyclization to obtain the plasmid pCENSK (Fig. 33). plasmid pBI121KS constructed in Example 9 was digested with the restriction enzymes KpnI and SalI to remove glucuronidase gene and, instead thereof, a DNA fragment

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containing the present CTP-CP4 EPSPS gene, which was obtained by digesting the above plasmid pCENSK with the restriction enzymes KpnI and SalI, was inserted therein to construct the plasmid pBICE (Fig. 34) wherein the present CTP-CP4 EPSPS gene was ligated to the downstream of 35S promoter.

The plasmid pBICE was introduced into Agrobacterium tumefaciens LBA44044 (manufactured by Clontech) and this was cultured in LB medium (0.5% yeast extract, 1.0% Bacto tryptone, 0.5% NaCl) containing 300 µg/ml of streptomycin, 100 µg/ml of rifampicin and 25 µg/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBICE.

Tobacco leaf pieces cultivated sterilely were infected with the Agrobacterium strain and, according to the same manner as in Example 5, tobacco bearing the inserted present CTP-CP4 EPSPS gene was obtained.

Example 48

Introduction of Present CTP-CP4 EPSPS Gene and Gene Encoding Variant Tobacco Chelatase Subunit into Tobacco

A plasmid was constructed for introducing the present CTP-CP4 EPSPS gene and a gene encoding a variant tobacco chelatase subunit into a plant by Agrobacterium infection method. First, the plasmid pCENSK constructed in

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Example 47 was digested with the restriction enzyme KpnI to obtain a DNA fragment containing the present CTP-CP4 EPSPS gene, a terminator of a gene encoding nopaline synthase located at the downstream thereof and 35S promoter located at the downstream of the terminator. This was inserted into the KpnI restriction site of the plasmid pBITCHLH constructed in Example 9 to construct the plasmid pBICETCH (Fig. 35) wherein the present CTP-CP4 EPSPS gene and the gene encoding the variant tobacco chelatase subunit were ligated to the downstream of 35S promoter, respectively.

The plasmid pBICETCH was introduced into Agrobacterium tumefaciens LBA44044 and this was cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBICETCH.

Tobacco leaf pieces cultivated sterilely were infected with the Agrobacterium strain and, according to the same manner as described in Example 5, tobacco bearing the inserted present CTP-CP4 EPSPS gene and gene encoding the variant tobacco chelatase subunit was obtained.

Example 49

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Present CTP-CP4
EPSPS Gene as well as Tobacco Bearing Introduced Present

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CTP-CP4 EPSPS Gene and Gene Encoding Variant Tobacco Chelatase Subunit

Leaves of the tobacco bearing the introduced present CTP-CP4 EPSPS gene produced in Example 47, those of the control recombinant tobacco obtained in Example 5 and those bearing the introduced present CTP-CP4 gene and gene encoding the variant tobacco chelatase subunit produced in Example 48 are collected, and each leaf is divided into the right and left equivalent pieces along the main vein. of the pieces is treated with an aqueous solution containing 0.3 ppm PPO inhibitory-type herbicidal compound of Structure 8, while to the other piece is not treated with the compound. These leaf pieces are placed on MS medium containing 0.8% agar and allowed to stand at room temperature for 7 days in a light place. Then, each leaf piece is ground in 5 ml of 80% aqueous acetone solution in a mortar with a pestle to extract chlorophyll. The extract is diluted 10 times with 80% aqueous acetone solution and the absorbance is measured at 750 nm, 663 nm and 645 nm to calculate the total chlorophyll content according to the method described by Macknney G., J. Biol. Chem. (1941) 140, The resistant level to the herbicidal compound tested is represented by the percentage of the total chlorophyll content of the leave piece treated with the herbicidal compound to that of untreated leaf piece.

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Similarly, the tobacco bearing the introduced present CTP-CP4 EPSPS gene, the tobacco bearing the introduced both present CTP-CP4 EPSPS gene and gene encoding the variant tobacco chelatase subunit, and the control recombinant tobacco are treated with an aqueous solution containing 100 ppm of a glyphosate to determine the resistant level to the glyphosate. The resistant level to the glyphosate is represented by the percentage of the total chlorophyll content of the leave piece treated with the glyphosate to that of untreated leaf piece.

Example 50

Introduction of Present CTP-CP4 EPSPS Gene and Gene Encoding Variant Soybean PPO into Tobacco

A plasmid was constructed for introducing the present CTP-CP4 EPSPS gene and a gene encoding a variant soybean PPO into a plant by Agrobacterium infection method. According to the same manner as described in Example 11, PCR was carried out by using an oligonucleotide primer composed of the nucleotide sequence represented by SEQ ID NO: 75, an oligonucleotide primer composed of the nucleotide sequence represented by SEQ ID NO: 76, and the plasmid pSPPO-P constructed in Example 3 as a template to amplify a DNA fragment containing the gene encoding the variant soybean PPO. Then, the plasmid pBI121K constructed in Example 9 was digested with restriction enzymes KpnI and

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2.0

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SacI to remove β -glucuronidase gene and, instead thereof, a DNA fragment obtained by digesting the above DNA fragment containing the gene encoding the variant soybean PPO with the restriction enzymes KpnI and SacI was inserted therein to construct the plasmid pBIGMP (Fig. 36) wherein the gene was ligated to the downstream of the 35S promoter.

Then, the plasmid pCENSK constructed in Example 47 is digested with the restriction enzyme KpnI to obtain a DNA fragment containing the present CTP-CP4 EPSPS gene, the terminator of the gene encoding nopaline synthase located at the downstream of the gene and the 35S promoter located at the downstream of the terminator, followed by insertion of it in the KpnI restriction site of the above plasmid pBIGMP to construct the plasmid pBICEGMP (Fig. 37) wherein the present CTP-CP4 EPSPS gene and the gene encoding the variant soybean PPO are ligated to the downstream of the 35S promoter, respectively.

The plasmid pBICEGMP is introduced into Agrobacterium tumefaciens LBA44044 and this is cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBICEGMP.

Tobacco leaf pieces cultivated sterilely are infected with the Agrobacterium strain and, according to

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the same manner as in Example 5, tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the variant soybean PPO is obtained.

Example 51

Confirmation of Resistance to Herbicidal Compound of Tobacco Bearing Introduced Present CTP-CP4 EPSPS Gene and Gene Encoding Variant Soybean PPO

The levels of resistance to the PPO inhibitory-type herbicidal compound represented by structure 8 are confirmed quantitatively by testing the tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the variant soybean PPO obtained in Example 50, and the control recombinant tobacco obtained in Example 5 according to the same manner as in Example 49.

Further the levels of resistance to glyphosate are confirmed quantitatively by testing the tobacco bearing introduced the present CTP-CP4 EPSPS gene and the gene encoding the variant soybean PPO and the control recombinant tobacco according to the same manner as in Example 49.

Example 52

Introduction of Present CTP-CP4 EPSPS Gene and Gene Encoding Variant Chlamydomonas reinhardtii PPO into Tobacco

A plasmid was constructed for introducing the

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present CTP-CP4 EPSPS gene and a gene encoding a variant Chlamydomonas reinhardtii PPO into a plant by Agrobacterium The plasmid pTVCRP constructed in infection method. Example 16 was digested with the restriction enzymes BamHI and SacI to prepare a DNA fragment containing a gene encoding a variant Chlamydomonas reinhardtii PPO. Binary vector pBI121 (manufactured by Clontech) was digested with the restriction enzymes BamHI and SacI to remove glucuronidase gene and, instead thereof, the above DNA encoding the containing the gene fragment Chlamydomonas reinhardtii PPO was inserted therein to construct the plasmid pBICRP (Fig. 38) wherein the gene was ligated to the downstream of the 35S promoter.

Then, the plasmid pBICRP is digested with the restriction enzyme BamHI and nucleotides are added to the gap of the double-stranded DNA with DNA Polymerase I to blunt the end of the DNA. The 5'-end of the DNA is dephosphorylated by treatment with an alkaline phosphatase derived from calf small intestine, followed by inserting a phosphorylated KpnI linker (4668A manufactured by Takara Shuzo Co., Ltd.) and cyclization to obtain plasmid pBICRPK. Then, the plasmid pCENSK constructed in Example 47 is digested with the restriction enzyme KpnI to obtain a DNA fragment containing the present CTP-CP4 EPSPS gene, the terminator of the gene encoding nopaline synthase located

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at the downstream of the present CTP-CP4 EPSPS gene and the 35S promoter located at the downstream of the terminator. This is inserted in the KpnI restriction site of the above plasmid pBICRPK to construct the plasmid pBICECRP (Fig. 39) wherein the present CTP-CP4 EPSPS gene and the gene encoding the variant Chlamydomonas reinhardtii PPO are ligated to the downstream of 35S promoter, respectively.

The plasmid pBICECRP is introduced into Agrobacterium tumefaciens LBA44044 and this is cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBICECRP.

Tobacco leaf pieces cultivated sterilely are infected with the Agrobacterium strain and, according to the same manner as described in Example 5, tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the variant Chlamydomonas reinhardtii PPO is obtained.

Example 53

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Present CTP-CP4
EPSPS Gene and Gene Encoding Variant Chlamydomonas
reinhardtii PPO

The levels of resistance to the above PPO inhibitory-type herbicidal compound represented by

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Structure 8 are confirmed quantitatively by testing the tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the variant *Chlamydomonas reinhardtii* PPO obtained in Example 52 and the control recombinant tobacco obtained in Example 5 according to the same manner as in Example 49.

Further the levels of resistance to glyphosate are confirmed quantitatively by testing the tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the variant *Chlamydomonas reinhardtii* PPO and the control recombinant tobacco according to the same manner as in Example 49.

Example 54

Introduction of Present CTP-CP4 EPSPS Gene and Gene Encoding Chloroplast-Localized Type Ferrochelatase of Arabidopsis thaliana into Tobacco

The plasmid pBIATF constructed in Example 39 is digested with the restriction enzyme BamHI and then nucleotides are added in the gap of the double-stranded DNA with DNA polymerase I to blunt the end of the DNA. The 5'-end of the DNA is dephosphorylated by treatment with an alkaline phosphatase derived from calf small intestine, followed by insertion of a phosphorylated KpnI linker (4668A manufactured by Takara Shuzo Co., Ltd.) and cyclization to obtain the plasmid pBIATFK. Then, the

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plasmid pCENSK constructed in Example 47 is digested with the restriction enzyme KpnI to obtain a DNA fragment containing the present CTP-CP4 EPSPS gene, the terminator of the gene encoding nopaline synthase located at the downstream of the present CTP-CP4 EPSPS gene and the 35S promoter located at the downstream of the terminator. This is inserted in the KpnI restriction site of the above plasmid pBIATFK to construct the plasmid pBICEATF (Fig. 40) wherein the present CTP-CP4 EPSPS gene and the gene encoding the chloroplast-localized type ferrochelatase of Arabidopsis thaliana are ligated to the downstream of 35S promoter, respectively.

The plasmid pBICEATF is introduced into Agrobacterium tumefaciens LBA44044 and this is cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBICEATF.

Tobacco leaf pieces cultivated sterilely are infected with the Agrobacterium strain and, according to the same manner as described in Example 5, tobacco bearing the inserted present CTP-CP4 EPSPs gene and gene encoding the chloroplast-localized type ferrochelatase of Arabidopsis thaliana is obtained.

Example 55

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Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Present CTP-CP4
EPSPS Gene and Gene Encoding Chloroplast-Localized Type
Ferrochelatase of Arabidopsis thaliana

The levels of resistance to the above PPO inhibitory-type herbicidal compound represented by Structure 8 are confirmed quantitatively by testing the tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the chloroplast-localized type ferrochelatase of Arabidopsis thaliana obtained in Example 54, and the control recombinant tobacco obtained in Example 5 according to the same manner as in Example 49.

Further the levels of resistance to glyphosate are confirmed quantitatively by testing the tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the chloroplast-localized type ferrochelatase of Arabidopsis thaliana and the control recombinant tobacco according to the same manner as in Example 49.

Example 56

Introduction of Present CTP-CP4 EPSPS Gene and Gene Encoding Present Soybean CPOX into Tobacco

A plasmid was constructed for introducing the present CTP-CP4 EPSPS gene and a gene encoding the present soybean CPOX into a plant by Agrobacterium infection method. First, the plasmid pCRSCPOX constructed in Example 42 was

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digested with the restriction enzyme BamHI to prepare a DNA fragment containing a gene encoding the present soybean CPOX. The DNA fragment was inserted in the BamHI restriction site of the plasmid pBI121KS constructed in Example 9 to obtain the plasmid pBISCPOXGUS. This plasmid was digested with the restriction enzyme SalI to remove β -glucuronidase gene, followed by self-cyclization to construct the plasmid pBISCPOX (Fig. 41) wherein the gene was ligated to the downstream of the 35S promoter.

Then, the plasmid pBISCPOX is digested with the restriction enzyme BamHI and nucleotides are added to the gap of the double-stranded DNA with DNA polymerase I to blunt the end of the DNA. The 5'-end of the DNA is dephosphorylated by treatment with an alkaline phosphatase derived from calf small intestine, followed by inserting a phosphorylated KpnI linker (4668A manufactured by Takara Shuzo Co., Ltd.) therein and cyclization to obtain the plasmid pBISCPOXK. Then, the plasmid pCENSK constructed in Example 47 is digested with the restriction enzyme KpnI to obtain a DNA fraction containing the present CTP-CP4 EPSPS gene, the terminator of the gene encoding nopaline synthase located at the downstream of the present CTP-CP4 EPSPS gene and the 35S promoter located at the downstream of the terminator. This is inserted in the KpnI restriction site of the above plasmid pBISCPOXK to construct the plasmid

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pBICESCPOX (Fig. 42) wherein the present CTP-CP4 EPSPS gene and the gene encoding the present soybean CPOX are ligated to the downstream of 35S promoter, respectively.

The plasmid pBICESCPOX is introduced into Agrobacterium tumefaciens LBA44044 and this is cultured in LB medium containing 300 μ g/ml of streptomycin, 100 μ g/ml of rifampicin and 25 μ g/ml of kanamycin, followed by selection of a transformant to isolate an Agrobacterium strain bearing pBICESCPOX.

Tobacco leaf pieces cultivated sterilely are infected with the *Agrobacterium* strain and, according to the same manner as described in Example 45, tobacco bearing the inserted present CTP-CP4 EPSPS gene and gene encoding the present soybean CPOX is obtained.

Example 57

Confirmation of Resistance to Herbicidal
Compounds of Tobacco Bearing Introduced Present CTP-CP4
EPSPS Gene and Gene Encoding Present Soybean CPOX

The levels of resistance to the above PPO inhibitory-type herbicidal compound represented by Structure 8 are confirmed quantitatively by testing the tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the present soybean CPOX obtained in Example 56, and the control recombinant tobacco according to the same manner as in Example 49.

Further the levels of resistance to glyphosate are confirmed quantitatively by testing the tobacco bearing the introduced present CTP-CP4 EPSPS gene and gene encoding the present soybean CPOX and the control recombinant tobacco according to the same manner as in Example 49.

As described hereinabove, according to the present invention, weed control compound-resistant plant can be produced.

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fragment having partial sequence of tobacco chlH gene SEQ ID NO: 11 Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene 5 SEO ID NO: 12 Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene SEO ID NO: 13 Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene 10 SEO ID NO: 14 Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene SEO ID NO: 15 amplify 15 Designed oligonucleotide primer to Chlamydomonas PPO gene SEO ID NO: 16 Designed oligonucleotide primer to amplify Chlamydomonas PPO gene SEO ID NO: 19 20 Designed oligonucleotide primer to amplify DNA fragment having partial sequence of Chlamydomonas PPO gene SEO ID NO: 20

Designed oligonucleotide primer to amplify DNA

fragment having partial sequence of Chlamydomonas PPO gene

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SEO ID NO: 21

Designed oligonucleotide primer to amplify DNA fragment having partial sequence of cucumber ferrochelatase gene

SEQ ID NO: 22

Designed oligonucleotide primer to amplify DNA fragment having partial sequence of cucumber ferrochelatase gene

SEQ ID NO: 23

Designed oligonucleotide primer to amplify $Escherichia \ coli \ hemF \ gene$

SEQ ID NO: 24

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SEQ ID NO: 25

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SEO ID NO: 26

Designed oligonucleotide primer to amplify

20 Escherichia coli hemF gene

SEO ID No: 27

Designed oligonucleotides to synthesize genes encoding random peptides comprising 5 amino acids

SEO ID NO: 28

Designed oligonucleotides to synthesize genes

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encoding random peptides comprising 5 amino acids

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Designed oligonucleotide to synthesize the gene encoding the peptide HASYS

SEO ID NO: 30

Designed oligonucleotide to synthesize the gene encoding the peptide HASYS

SEQ ID NO: 31

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SEQ ID NO: 32

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Designed oligonucleotide to synthesize the gene encoding the peptide MGHASYS

SEQ ID NO: 34

Designed oligonucleotide to synthesize the gene encoding the peptide MGHASYS $\begin{tabular}{ll} \end{tabular} \label{eq:model}$

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SEO ID NO: 38

Designed oligonucleotide to synthesize the gene encoding the peptide MGYAGY

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Designed oligonucleotide to synthesize the gene encoding the peptide $\ensuremath{\mathsf{MGYAGF}}$

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Designed oligonucleotide to synthesize the gene encoding the peptide ${\tt MGYAGF}$

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20 Designed oligonucleotide to synthesize the gene encoding the peptide MG(HASYS)4

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SEQ ID NO: 51

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encoding the peptide MG(RASSL)8

SEQ ID NO: 52

Designed oligonucleotide to synthesize the gene encoding the peptide MG(RASSL)8 $\,$

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Protoporphyrin IX binding protein HASYS

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SEO ID NO: 54

Protoporphyrin IX binding protein MGHASYS

SEQ ID NO: 55

Protoporphyrin IX binding protein RASSL

SEQ ID NO: 56

Protoporphyrin IX binding protein MGRASSL

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H2TMpyP binding protein MGYAGY

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H2TMpyP binding protein YAGF

SEQ ID NO: 60

H2TMpyP binding protein MGYAGF

SEQ ID NO: 61

Protoporphyrin IX binding protein MG(HASYS)

SEQ ID NO: 62

Protoporphyrin IX binding protein MG(HASYS)

SEO ID NO: 63

Protoporphyrin IX binding protein MG(RASSL),

SEQ ID NO: 64

Protoporphyrin IX binding protein MG(RASSL).

SEO ID NO: 65

Designed oligonucleotide primer to introduce

25 mutation into arabidopsis PPO gene

SEO ID NO: 66

 $\label{eq:Designed} Designed \quad \text{oligonucleotide} \quad primer \quad \text{to} \quad amplify \\ herbicide \quad resistant \quad arabidopsis \quad PPO(A220V) \quad gene$

SEO ID NO: 67

Designed oligonucleotide primer to amplify herbicide resistant arabidopsis PPO(A220V) gene

SEQ ID NO: 68

Designed oligonucleotide primer to amplify arabidopsis chloroplast ferrochelatase gene

SEQ ID NO: 69

Designed oligonucleotide primer to amplify arabidopsis chloroplast ferrochelatase gene

SEQ ID NO: 70

Designed oligonucleotide primer to amplify soybean coproporphyrinogen III oxidase gene

SEQ ID NO: 71

 $\label{eq:decomposition} \mbox{Designed oligonucleotide primer to amplify} \\ \mbox{soybean coproporphyrinogen III oxidase gene}$

SEQ ID NO: 72

Designed oligonucleotide primer to amplify DNA fragment having nucleotide sequence encoding the Petunia hybrida EPSPS chloroplast transit peptide and the Agrobacterium sp. strain CP4 EPSPS gene

SEQ ID NO: 73

Designed oligonucleotide primer to amplify DNA

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fragment having nucleotide sequence encoding the Petunia hybrida EPSPS chloroplast transit peptide and the Agrobacterium sp. strain CP4 EPSPS gene

SEO ID NO: 75

5 Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene
SEO ID NO: 76

Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene

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<211> 35

<212> DNA

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<220>

 $\ensuremath{^{<\!223>}}$ Designed oligonucleotide primer to amplify DNA fragment having

partial sequence of tobacco chlH gene

<400> 9

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20 <210> 10

<211> 34

<212> DNA

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<211> 39

<212> DNA

<213> Artificial Sequence

<220>

<223> Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene

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<210> 12

<211> 36

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partial sequence of soybean PPO gene

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partial sequence of soybean PPO gene
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12207

 $\langle 223 \rangle$ Designed oligonucleotide primer to amplify Chlamydomonas PPO

gene

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10

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30

45

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20

5

25

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40

142

190

94

gcg acc gcg gcg gcc cgc cgc aca ctc cac cgc act gct gcg gcg gcc Ala Thr Ala Ala Ala Arg Arg Thr Leu His Arg Thr Ala Ala Ala Ala

•

25

20

50

35

55

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	Phe	Asp	Tyr	Pro	Pro	Val	G1y	Ala	Val	Thr	Leu	Ser	Tyr	Pro	Leu	Ser	
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	gcc	gtg	cgg	gag	gag	cgc	aag	gcc	tcg	gac	ggg	tcc	gtg	ccg	ggc	ttc	1246
	Ala	Val	Arg	G1u	Glu	Arg	Lys	Ala	Ser	Asp	Gly	Ser	Val	Pro	G1y	Phe	
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	Tyr	Ser	Ser	Ser	Leu	Phe	Pro	G1y	Arg	Ala	Pro	G1u	Gly			Leu	
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ctg	ctc	aac	tac	atc	ggc	ggc	acc	acc	aac	cgc	ggc	atc	gtc	aac	cag	1390
Leu	Leu	Asn	Tyr	Ile	G1y	Gly	Thr	Thr	Asn	Arg	Gly	Ile	Val	Asn	G1n	
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Thr	Thr	Glu	G1n	Leu	Val	G1u	G1n	Val	Asp	Lys	Asp	Leu	Arg	Asn	Met	
	465					470					475		,			
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Val	Ile	Lys	Pro	Asp	Ala	Pro	Lys	Pro	Arg	Val	Val	Gly	Val	Arg	Va1	
480					485					490					495	
tgg	ccg	cgc	gcc	atc	ccg	cag	ttc	aac	ctg	ggc	cac	ctg	gag	cag	ctg	1534
Trp	Pro	Arg	Ala	Ile	Pro	Gln	Phe	Asn	Leu	Gly	His	Leu	Glu	Gln	Leu	
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Asp	Lys	Ala	Arg	Lys	Ala	Leu	Asp	Ala	Ala	Gly	Leu	G1n	Gly	Val	His	
			515					520					525			
ctg	ggg	ggc	aac	tac	gtc	agc	ggt	gtg	gcc	ctg	ggc	aag	gtg	gtg	gag	1630
Leu	G1 y	G1y	Asn	Tyr	Val	Ser	Gly	Val	Ala	Leu	Gly	Lys	Val	Val	G1u	
		530					535					540				
cac	ggc	tac	gag	tcc	gca	gcc	aac	ctg	gcc	aag	agc	gtg	tcc	aag	gcc	1678
His	Gly	Tyr	G1u	Ser	Ala	Ala	Asn	Leu	Ala	Lys	Ser	Val	Ser	Lys	Ala	
	545					550					555					
gca	gtc	aag	gcc	taa	gcg	gctg	cag	cagt	agca	gc a	gcag	catc	g gg	ctgt	agct	1733
Ala	Va1	Lys	Ala													
560			563													
aat	anat	acc	acaa	taac	90 0	aaca	ocao	า ลล	ttøø	caag	cac	ttgg	ggc	aagc	ggagtg	1793

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1838

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<211> 563

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<213> Chlamydomonas reinhardtii CC407

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Arg Pro Thr Pro Phe Ser Val Ala Ser Pro Ala Thr Ala Ala Ser Pro
35 40 45

Ala Thr Ala Ala Ala Arg Arg Thr Leu His Arg Thr Ala Ala Ala Ala 50 55 60

Thr Gly Ala Pro Thr Ala Ser Gly Ala Gly Val Ala Lys Thr Leu Asp
65 70 75

Asn Val Tyr Asp Val Ile Val Val Gly Gly Gly Leu Ser Gly Leu Val

Thr Gly Gln Ala Leu Ala Ala Gln His Lys Ile Gln Asn Phe Leu Val

Thr Glu Ala Arg Glu Arg Val Gly Gly Asn Ile Thr Ser Met Ser Gly

115 120 125

	Asp	G1y	Tyr	Val	Trp	Glu	Glu	G1y	Pro	Asn	Ser	Phe	G1n	Pro	Asn	Asp
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	160					165					170			,		175
	Arg	Pro	Va1	Pro	Ser	G1y	Leu	Asp	Ala	Phe	Thr	Phe	Asp	Leu	Met	Ser
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	Ile	Pro	G1y	Lys	Ile	Arg	Ala	Gl y	Leu	Gly	Ala	He	G1y	Leu	Пе	Asn
.0				195					200					205		
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	240					245					250					255
	Asn	Arg	Ile	Trp	Ile	Leu	Glu	Lys	Asn	G1y	G1y	Ser	Leu	Val	G1y	G1y
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	Ala	Ile	Lys	Leu	Phe	G1n	Glu	Arg	Gln	Ser	Asn	Pro	Ala	Pro	Pro	Arg
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			290					295					300			
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		Asp	G1y	Arg	Tyr	G1y	Leu	Val	Tyr	Asp	Thr	Pro	Glu	G1 y	Arg	Val	Lys
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	5	Va1	Phe	Ala	Arg	Ala	Val	Ala	Leu	Thr	Ala	Pro	Ser	Tyr	Val	Val	Ala
					355					360					365		
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II. Some there thank staff Sault Sault				370					375					380			
		Phe	Asp	Tyr	Pro	Pro	Va1	G1y	Ala	Val	Thr	Leu	Ser	Tyr	Pro	Leu	Ser
	10		385					390					395				
		Ala	Val	Arg	Glu	G1u	Arg	Lys	Ala	Ser	Asp	G1y	Ser	Val	Pro	G1y	Phe
Geerl		400					405					410					415
though these firms there thank at		G1y	Gln	Leu	His	Pro	Arg	Thr	G1n	G1y	Ile	Thr	Thr	Leu	G1y	Thr	Ile
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	20		465					470					475				
		Val	Ile	Lys	Pro	Asp	Ala	Pro	Lys	Pro	Arg	Va1	Val	Gly	Val	Arg	Val
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<210> 21

10 <211> 33

<212> DNA

<213> Artificial Sequence

<220>

<223> Designed oligonucleotide primer to amplify DNA fragment having partial sequence of cucumber ferrochelatase gene

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<211> 36

<212> DNA

<213> Artificial Sequence

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<210> 23

<211> 25

10 <212> DNA

<213> Artificial Sequence

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<223> Designed oligonucleotide primer to amplify Esherichia coli hemF

15 gene

<400> 23

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20 <210> 24

<211> 24

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25
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10 peptide MGHASYS

<400> 33

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15 <210> 34

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<213> Artificial Sequence

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 $\langle 223 \rangle$ Designed oligonucleotide to synthesize the gene encoding the peptide MGYAGY

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<210> 38

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<213> Artificial Sequence

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5 $\langle 223 \rangle$ Designed oligonucleotide to synthesize the gene encoding the peptide MG(HASYS)4

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<210> 42

⟨211⟩ 34

<212> DNA

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<210> 43

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25 <212> DNA

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<400> 43

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<212> DNA

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<210> 45

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<212> DNA

25 <213> Artificial Sequence

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<400> 45

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<210> 47

<211> 34

<212> DNA

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<223> Designed oligonucleotide to synthesize the gene encoding the peptide MG(RASSL)4

5 <400> 47

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10 <212> DNA

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20 <210> 49

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5 <210> 54

(211) 7

<212> PRT

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15 1

5

<210> 55

⟨211⟩ 5

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25 <400> 55

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<400> 57
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Ser His Ala Ser Tyr Ser
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,Ō		1 5 10 15
		Ser His Ala Ser Tyr Ser His Ala Ser Tyr Ser His Ala Ser Tyr Ser
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	15	His Ala Ser Tyr Ser His Ala Ser Tyr Ser
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<223> Designed oligonucleotide primer to introduce mutation into arabidopsis PPO gene

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10 <210> 66

<211> 32

<212> DNA

<213> Artificial Sequence

15 <220>

 $\langle 223 \rangle$ Designed oligonucleotide primer to amplify herbicide resistant arabidopsis PPO(A220V) gene

<400> 66

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<210> 67

<211> 34

<212> DNA

25 <213> Artificial Sequence

<213> Designed oligonucleotide primer to amplify herbicide resistant arabidopsis PPO(A220V) gene

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<400> 67

gggagattta atgtcgacca tttacttgta agcg 34

<210> 68

<211> 31 10

<212> DNA

<213> Artificial Sequence

<220>

amplify Arabidopsis 15 <223> Designed oligonucleotide primer to chloroplast ferrochelatase gene

<400> 68

31 gateggttet gaaatttgga teeatgeagg e

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<210> 69

<211> 31

<212> DNA

<213> Artificial Sequence

<223> Designed oligonucleotide primer to amplify Arabidopsis chloroplast ferrochelatase gene

5 <400> 69

cacaaaacca acgagctcct ataggttccg g 31

<210> 70

<211> 30

10 <212> DNA

<213> Artificial Sequence

<220>

 $\langle 223 \rangle$ Designed oligonucleotide primer to amplify soybean

15 coproporphyrinogen III oxidase gene

<400> 70

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20 <210> 71

<211> 30

<212> DNA

<213> Artificial Sequence

25 <220>

<223> Designed oligonucleotide primer to amplify soybean coproporphyrinogen III oxidase gene

<400> 71

5 gggggtcgac tgatgaatta gatccattcc 30

<210> 72

<211> 36

<212> DNA

10 <213> Artificial Sequence

<220>

<223> Designed oligonucleotide primer to amplify DNA fragment having nucleotide sequence encoding the Petunia hybrida EPSPS chloroplast transit peptide and the Agrobacterium sp. strain CP4 EPSPS gene

<400> 72

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20 <210> 73

<211> 32

<212> DNA

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25 <220>

<223>Designed oligonucleotide primer to amplify DNA fragment having nucleotide sequence encoding the Petunia hybrida EPSPS chloroplast transit peptide and the Agrobacterium sp. strain CP4 EPSPS gene

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10 <212> DNA

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<220>

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aat tcc aat ttc cat aaa ccc caa gtt cct aaa tct tca agt ttt ctt 96
Asn Ser Asn Phe His Lys Pro Gln Val Pro Lys Ser Ser Ser Phe Leu

20 25 30

	gtt	ttt	gga	tct	aaa	aaa	ctg	aaa	aat	tca	gca	aat	tct	atg	ttg	gtt	144
	Val	Phe	G1y	Ser	Lys	Lys	Leu	Lys	Asn	Ser	Ala	Asn	Ser	Met	Leu	Val	
				35					40					45			
	ttg	aaa	aaa	gat	tca	att	ttt	atg	caa	aag	ttt	tgt	tcc	ttt	agg	att	192
5	Leu	Lys	Lys	Asp	Ser	I1e	Phe	Met	G1n	Lys	Phe	Cys	Ser	Phe	Arg	Ile	
			50					55					60				
	tca	gca	tca	gtg	gct	aca	gcc	$_{ m tgc}$	atg	ctt	cac	ggt	gca	agc	agc	cgg	240
	Ser	Ala	Ser	Val	Ala	Thr	Ala	Cys	Met	Leu	His	Gly	Ala	Ser	Ser	Arg	
		65					70					75					
10	ccc	gca	acc	gcc	cgc	aaa	tcc	tct	ggc	ctt	tcc	gga	acc	gtc	cgc	att	288
	Pro	Ala	Thr	Ala	Arg	Lys	Ser	Ser	Gly	Leu	Ser	Gly	Thr	Val	Arg	Ile	
	80					85					90					95	
	ccc	ggc	gac	aag	tcg	atc	tcc	cac	cgg	tcc	ttc	atg	ttc	ggc	ggt	ctc	336
	Pro	G1y	Asp	Lys	Ser	Ile	Ser	His	Arg	Ser	Phe	Met	Phe	Gly	Gly	Leu	
15					100					105					110		
	gcg	agc	ggt	gaa	acg	cgc	atc	acc	ggc	ctt	ctg	gaa	ggc	gag	gac	gtc	384
	Ala	Ser	Gly	Glu	Thr	Arg	Ile	Thr	G1y	Leu	Leu	Glu	Gly	Glu	Asp	Val	
				115					120					125			
	atc	aat	acg	ggc	aag	gcc	atg	cag	gcc	atg	ggc	gcc	agg	atc	cgt	aag	432
20	I1e	Asn	Thr	Gly	Lys	Ala	Met	G1n	Ala	Met	G1y	Ala	Arg	Ile	Arg	Lys	
			130					135					140				
	gaa	ggc	gac	acc	tgg	atc	atc	gat	ggc	gtc	ggc	aat	ggc	ggc	ctc	ctg	480
	G1u	Gly	Asp	Thr	Trp	Ile	Ile	Asp	G1y	Val	Gly	Asn	G1y	Gly	Leu	Leu	
		145					150					155					

	gcg	cct	gag	gcg	ccg	ctc	gat	ttc	ggc	aat	gcc	gcc	acg	ggc	tgc	cgc	528
	Ala	Pro	Glu	Ala	Pro	Leu	Asp	Phe	G1y	Asn	Ala	Ala	Thr	G1y	Cys	Arg	
	160					165					170					175	
	ctg	acc	atg	ggc	ctc	gtc	ggg	gtc	tac	gat	ttc	gac	agc	acc	ttc	atc	576
5	Leu	Thr	Met	Gly	Leu	Val	Gly	Val	Tyr	Asp	Phe	Asp	Ser	Thr	Phe	Ile	
					180					185					190		
	ggc	gac	gcc	tcg	ctc	aca	aag	cgc	ccg	atg	ggc	cgc	gtg	ttg	aac	ccg	624
	Gly	sp /	Ala S	Ser l	Leu î	Γhr I	_ys /	Arg l	Pro l	Met (Gly A	Arg '	Val 1	Leu <i>i</i>	Asn l	Pro	
				195					200					205			
10	ctg	cgc	gaa	atg	ggc	gtg	cag	gtg	aaa	tcg	gaa	gac	ggt	gac	cgt	ctt	672
	Leu	Arg	G1u	Met	Gly	Val	G1n	Val	Lys	Ser	G1u	Asp	G1y	Asp	Arg	Leu	
			210					215					220				
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	Pro	Val	Thr	Leu	Arg	G1y	Pro	Lys	Thr	Pro	Thr	Pro	Ile	Thr	Tyr	Arg	
15		225					230					235					
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	Val	Pro	Met	Ala	Ser	Ala	G1n	Val	Lys	Ser	Ala	Val	Leu	Leu	Ala	G1y	
	240					245					250					255	
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20	Leu	Asn	Thr	Pro	Gly	Ile	Thr	Thr	Val	Ile	Glu	Pro	Ile	Met	Thr	Arg	
					260					265					270		
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	Asp	His	Thr	G1u	Lys	Met	Leu	Gln	G1y	Phe	G1y	Ala	Asn	Leu	Thr	Val	
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		290					295					300				
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Lys	Leu	Thr	G1y	G1n	Val	Ile	Asp	Val	Pro	Gly	Asp	Pro	Ser	Ser	Thr	
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Ala	Phe	Pro	Leu	Val	Ala	Ala	Leu	Leu	Val	Pro	G1y	Ser	Asp	Val	Thr	
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Ile	Leu	Asn	Val	Leu	Met	Asn	Pro	Thr	Arg	Thr	G1y	Leu	Ile	Leu	Thr	
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Leu	Gln	G1u	Met	Gly	Ala	Asp	Ile	G1u	Val	Ile	Asn	Pro	Arg	Leu	Ala	
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G1y	G1y	G1u	Asp	Val	Ala	Asp	Leu	Arg	Val	Arg	Ser	Ser	Thr	Leu	Lys	
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G1y	Val	Thr	Val	Pro	Glu	Asp	Arg	Ala	Pro	Ser	Met	Ile	Asp	Glu	Tyr	
	385					390					395					
ccg	att	ctc	gct	gtc	gcc	gcc	gcc	ttc	gcg	gaa	ggg	gcg	acc	gtg	atg	1248
Pro	Ile	Leu	Ala	Va1	Ala	Ala	Ala	Phe	Ala	Glu	Gly	Ala	Thr	Val	Met	
400					405					410					415	

	aac	ggt	ctg	gaa	gaa	ctc	cgc	gtc	aag	gaa	agc	gac	cgc	ctc	tcg	gcc	1296
	Asn	G1y	Leu	G1u	G1u	Leu	Arg	Val	Lys	Glu	Ser	Asp	Arg	Leu	Ser	Ala	
					420					425					430		
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5	Val	Ala	Asn	Gly	Leu	Lys	Leu	Asn	G1y	Va1	Asp	Cys	Asp	G1u	G1y	Glu	
				435					440					445			
	acg	tcg	ctc	gtc	gtg	cgc	ggc	cgc	cct	gac	ggc	aag	ggg	ctc	ggc	aac	1392
	Thr	Ser	Leu	Val	Val	Arg	Gly	Arg	Pro	Asp	G1y	Lys	G1y	Leu	Gly	Asn	
			450					455					460				
10	gcc	tcg	ggc	gcc	gcc	gtc	gcc	acc	cat	ctc	gat	cac	cgc	atc	gcc	atg	1440
	Ala	Ser	G1y	Ala	Ala	Va1	Ala	Thr	His	Leu	Asp	His	Arg	I1e	Ala	Met	
		46 5					470					475					
	agc	ttc	ctc	gtc	atg	ggc	ctc	gtg	tcg	gaa	aac	cct	gtc	acg	gtg	gac	1488
	Ser	Phe	Leu	Val	Met	G1 y	Leu	Val	Ser	Glu	Asn	Pro	Val	Thr	Val	Asp	
15	480					485					490					49 5	
	gat	gcc	acg	atg	atc	gcc	acg	agc	ttc	ccg	gag	ttc	atg	gac	ctg	atg	1536
	Asp	Ala	Thr	Met	I1e	Ala	Thr	Ser	Phe	Pro	G1u	Phe	Met	Asp	Leu	Met	
					500					505					510		
	gcc	ggg	ctg	ggc	gcg	aag	atc	gaa	ctc	tcc	gat	acg	aag	gct	gcc	tga	1584
20	Ala	G1y	Leu	G1y	Ala	Lys	Ile	Glu	Leu	Ser	Asp	Thr	Lys	Ala	Ala		
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	tga																1587

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25 <211> 33

<212> DNA

<213> Artificial Sequence

<220>

5 <223> Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene

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<210> 76

<211> 33

<212> DNA

<213> Artificial Sequence

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<220>

<400> 76

<223> Designed oligonucleotide primer to amplify DNA fragment having partial sequence of soybean PPO gene

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cctgcagetc gagagetect actatttgta cac 33

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What is claimed is:

 A method for giving resistance to weed control compounds to plants which comprises the steps of:

introducing a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- $\begin{tabular}{ll} (c) & being & substantially & free & from & framework \\ regions & of variable & regions & in an immunoglobulin, \\ into & a plant cell; and \\ \end{tabular}$

expressing the gene.

- 2. The method according to claim 1, wherein the gene is introduced into the plant cell in the form that it is operably ligated to a promoter and a terminator both of which are functional in the plant cell.
- 3. The method according to claim 1 or 2, wherein the substance which is concerned with the weed control activity of the weed control compound is the weed control compound itself.
- 4. The method according to claim 1, wherein the substance which is concerned with the weed control activity

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of a weed control compound is an endogenous substance in a plant.

- 5. The method according to claim 1, wherein the weed control compound is that inhibiting porphyrin biosynthesis of a plant.
 - 6. The method according to claim 1, wherein the weed control compound is a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound.
- 7. The method according to claim 5 or 6, wherein the substance which is concerned with the weed control activity of the weed control compound is protoporphyrin IX.
- 8. The method according to claim 5 or 6, wherein the protein is protoporphyrin IX binding subunit protein of magnesium chelatase, or a variant of said protein having a specific affinity for protoporphyrin IX.
- 9. The method according to claim 8, wherein the protein is magnesium chelatase derived from a photosynthetic microorganism.
- 10. The method according to claim 8, wherein the protein is magnesium chelatase derived from a plant.
 - 11. The method according to claim 8, wherein the protein is magnesium chelatase derived from tobacco.
- 12. The method according to claim 5 or 6, wherein the protein comprises the amino acid sequence of SEQ

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TD NO: 53.

- 13. The method according to claim 5 or 6, wherein the protein has the amino acid sequence of SEQ ID NO: 54.
- 14. The method according to claim 5 or 6, wherein the protein comprises the amino acid sequence of SEQ ID NO: 55.
- \$15\$. The method according to claim 5 or 6, $wherein the protein has the amino acid sequence of SEQ ID <math display="inline">_{\mbox{\footnotesize{NO:}}}$ 56.
- 16. The method according to claim 5 or 6, wherein the protein comprises the amino acid sequence of SEQ ID NO: 57.
- 17. The method according to claim 5 or 6, wherein the protein has the amino acid sequence of SEQ ID NO: 58.
- 18. The method according to claim 5 or 6, wherein the protein comprises the amino acid sequence of SEQ $\scriptstyle\rm ID$ No: 59.
- 19. The method according to claim 5 or 6, wherein the protein has of the amino acid sequence of SEQ ID NO: 60.
 - \$20.\$ The method according to claim 5 or 6, wherein the protein is composed of 4 to 100 amino acids.
- 25 21. The method according to claim 5 or 6,

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wherein the substance which is concerned with the weed control activity of the weed control compound is protoporphyrinogen IX.

- 22. The method according to claim 5 or 6, wherein the protein is a variant of protoporphyrinogen IX oxidase having no capability of oxidizing protoporphyrinogen IX and having a specific affinity for a protoporphyrinogen IX.
- 23. The method according to claim 5 or 6, wherein the protein is a variant of protoporphyrinogen IX oxidase having no capability of oxidizing protoporphyrinogen IX and having a specific affinity for a protoporphyrin IX oxidase inhibitory-type herbicidal compound.
- 24. The method according to claim 22 or 23, wherein the protein is a variant of protoporphyrinogen IX oxidase derived from a plant.
- \$25.\$ The method according to claim 22 or 23, wherein the protein is a variant of protoporphyrinogen IX oxidase derived from soybean.
- 26. The method according to claim 22 or 23, wherein the protein is a variant of protoporphyrinogen IX oxidase derived from an algae.
- 27. The method according to claim 22 or 23, wherein the protein is a variant of protoporphyrinogen IX

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oxidase derived from Chlamydomonas.

28. A method for giving resistance to weed control compounds to plants which comprises the steps of:

introducing a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for protoporphyrin IX,
- (b) having substantially no capability of modifying protoporphyrinogen IX, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin, into a plant cell; and

expressing the gene.

- 29. The method according to claim 28, wherein the gene is introduced in the plant cell in the form that it is operably ligated to a promoter and a terminator both of which are functional in the plant cell.
- 30. The method according to claim 28, wherein the weed control compound is that inhibiting porphyrin biosynthesis of a plant.
- 31. The method according to claim 28, wherein the weed control compound is a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound.
- 32. The method according to claim 30 or 31, wherein the protein is magnesium chelatase or a variant of

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said protein having a specific affinity for protoporphyrin TX .

- 33. The method according to claim 30 or 31, wherein the protein is ferrochelatase or a variant of said protein having a specific affinity for protoporphyrin IX.
- 34. The method according to claim 30 or 31, wherein the protein is ferrochelatase derived from a plant.
- 35. The method according to claim 30 or 31, wherein the protein is ferrochelatase derived from barley.
- $\,$ 36. The method according to claim 30 or 31, wherein the protein is ferrochelatase derived from cucumber.
- $\,$ 37. The method according to claim 30 or 31, wherein the protein is a peptide composed of 4 to 100 amino acids.
- 38. A method for giving resistance to weed control compounds to plants which comprises the steps of:

introducing a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for protoporphyrinogen IX,
- (b) having the capability for modifying coproporphyrinogen III, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin;
- 25 into a plant cell; and

expressing the gene.

- 39. The method according to claim 38, wherein the gene is introduced into the plant cell in the form that it is operably ligated to a promoter and a terminator both of which are functional in the plant cell.
- 40. The method according to claim 38, wherein the protein is coproporphyrinogen III oxidase or a variant of said protein having a specific affinity for protoporphyrinogen IX.
- 41. The method according to claim 38, wherein the protein is coproporphyrinogen III oxidase derived from a microorganism.
- 42 The method according to claim 38, wherein the protein is coproporphyrinogen III oxidase derived from Escherichia coli.
- 43. A weed control compound-resistant plant whose resistance is given by the method of claim 1 or 28.
- 44. A weed control compound-resistant plant whose resistance is given by the method of claim 38.
- 45. A method for protecting a plant which comprises applying the weed control compound to a growth area of the plant of claim 43.
- 46. A method for protecting a plant which comprises applying said weed control compound to a growth area of the plant of claim 44.

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- 47. A method for selecting a plant which comprises applying a weed control compound to which the plant of claim 43 is resistant to a growth area of the plant of claim 43 and other plants, and selecting either plant on the basis of difference in growth between the plants.
- 48. A method for selecting a plant which comprises applying a weed control compound to which the plant of claim 44 is resistant to a growth area of the plant of claim 44 and other plants, and selecting either plant on the basis of difference in growth between the plants.
- \$49.\$ The method according to claim 47, wherein the plants are plant cells.
- 50. The method according to claim 48, wherein the plants are plant cells.
- wherein the weed control compound is a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound selected from the compounds of (1) to (3) below, and the substance which is concerned with the weed control activity of the weed control compound is protoporphyrin IX, protoporphyrinogen IX or a protoporphyrinogen IX oxidase inhibitory-type herbicidal compound:
 - (1) chlormethoxynil, bifenox, chlornitrofen,

acifluorfen and its ethyl ester, acifluorfen-sodium, oxyfluorfen, oxadiazon, 2-[4-chloro-2-fluoro-5-(prop-2-ynyloxy)phenyl]-2,3,4,5,6,7-hexahydro-1H-isoindol-1,3-dione, chlorphthalim, TNPP-ethyl, or N3-(1-phenylethyl)-2,6-dimethyl-5-propyonylnicotinamide;

(2) a compound represented by the general formula: J-G (I), wherein G is a group represented by any one of the following general formulas G-1 to G-9 and J is a group represented by any one of the following general formulas J-1 to J-30:

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$$R^{14}$$
 R^{14}
 R^{14}
 R^{14}
 R^{14}
 R^{14}
 R^{14}
 R^{15}
 R

wherein the dotted lines in the formulas J-5, J-6, J-12 and J-24 represent that the left hand ring contains only single bonds, or one bond in the ring is a double bond between carbon atoms;

X is oxygen atom or sulfur atom;

Y is oxygen atom or sulfur atom;

R1 is hydrogen atom or halogen atom;

 R^2 is hydrogen atom, C_1-C_8 alkyl group, C_1-C_8 haloalkyl group, halogen atom, OH group, OR^{27} group, SH group, $S(O)_pR^{27}$ group, COR^{27} group, CO_2R^{27} group,

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substituted with one or more and the same or different C_1 - C_4 alkyl groups;

p is 0, 1 or 2;

 $$R^3$$ is C_1-C_2 alkyl group, C_1-C_2 haloalkyl group, $OCH_3 \ group, \ SCH_3 \ group, \ OCHF_2 \ group, \ halogen \ atom, \ cyano group or nitro group;$

 R^4 is hydrogen atom, C_1-C_3 alkyl group, C_1-C_3 haloalkyl group or halogen atom;

 $\rm R^5$ is hydrogen atom, $\rm C_1-C_3$ alkyl group, halogen atom, $\rm C_1-C_3$ haloalkyl group, cyclopropyl group, vinyl group, $\rm C_2$ alkynyl group, cyano group, $\rm C(O)\,R^{38}$ group, $\rm CO_2R^{16}$ group, $\rm C(O)\,NR^{38}R^{39}$ group, $\rm CR^{34}R^{35}CO$ group, $\rm CR^{24}R^{35}CO_2R^{38}$ group, $\rm CR^{34}R^{35}CO_2R^{38}$ group, $\rm CR^{34}R^{35}CO_3R^{38}$ group, $\rm CR^{34}R^{35}CO_3R^{38}$ group, $\rm CR^{34}R^{35}CO_3R^{38}$ group, or, when G is G-2 or G-6, R⁴ and R⁵ may form C=O group together with the carbon atom to which they are attached;

 R^6 is C_1-C_6 alkyl group, C_1-C_6 haloalkyl group, C_2-C_6 alkoxyalkyl group, C_3-C_6 alkenyl group or C_3-C_6 alkynyl group;

 X^1 is single bond, oxygen atom, sulfur atom, NH group, $N(C_1-C_3$ alkyl) group, $N(C_1-C_3$ haloalkyl) group or N(allyl) group;

 R^7 is hydrogen atom, $C_1\text{--}C_6$ alkyl group, $C_1\text{--}C_6$ haloalkyl group, halogen atom, $S\left(0\right)_2\left(C_1\text{--}C_6\text{alkyl}\right)$ group or $C\left(=0\right)R^{40}$ group;

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 R^{8} is hydrogen atom, $C_{1}-C_{8}$ alkyl group, $C_{3}-C_{8}$ cycloalkyl group, $C_{3}-C_{8}$ alkenyl group, $C_{3}-C_{8}$ alkoxyalkyl group, $C_{1}-C_{8}$ haloalkyl group, $C_{2}-C_{8}$ alkoxyalkyl group, $C_{3}-C_{8}$ alkoxyalkyl group, $C_{3}-C_{8}$ alkoxyalkoxyalkyl group, $C_{3}-C_{8}$ haloalkynyl group, $C_{3}-C_{8}$ haloalkenyl group, $C_{1}-C_{8}$ alkylsulfonyl group, $C_{1}-C_{8}$ haloalkylsulfonyl group, $C_{1}-C_{8}$ alkoxycarbonylalkyl group, $S(O)_{2}NH(C_{1}-C_{8}$ alkyl) group, $C(O)_{2}NH(C_{1}-C_{8}$ alkyl) group, $C(O)_{2}NH(C_{1}-C_{8}$ alkyl) group, $C(O)_{2}NH(C_{1}-C_{8})$ group or benzyl group whose phenyl ring may be substituted with R^{42} ;

 $\ensuremath{\text{n}}$ and $\ensuremath{\text{m}}$ are independently 0, 1, 2 or 3 and $\ensuremath{\text{m}}$ + n is 2 or 3;

 $\label{eq:Z} Z \ \mbox{is CR^9R^{10} group, oxygen atom, sulfur atom, $S(0)$ group, $S(0)_2$ group or $N(C_1-C_4$ alkyl)$ group;}$

each R^9 is independently hydrogen atom, C_1-C_3 alkyl group, halogen atom, hydroxyl group, C_1-C_6 alkoxy group, C_1-C_6 haloalkyl group, C_1-C_6 haloalkyl group, C_2-C_6 alkylcarbonyloxy group or C_2-C_6 haloalkylcarbonyloxy group;

each R^{10} is independently hydrogen atom, $C_1 - C_3$ alkyl group, and hydroxyl group or halogen atom;

 R^{11} and R^{12} are independently hydrogen atom, halogen atom, C_1-C_6 alkyl group, C_3-C_6 alkenyl group or C_1-C_6 haloalkyl group;

 R^{13} is hydrogen atom, C_1-C_6 alkyl group, C_1-C_6 haloalkyl group, C_3-C_6 haloalkyl group, C_3-C_6 haloalkenyl group, C_3-C_6 haloalkynyl group

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 $$R^{14}$$ is $C_1^-C_6$ alkyl group, $C_1^-C_6$ alkylthio group, $C_1^-C_6$ haloalkyl group or $N\left(CH_3\right)_2$ group;

W is nitrogen atom or CR15;

 R^{15} is hydrogen atom, C_1-C_6 alkyl group, halogen atom, or phenyl group optionally substituted with C_1-C_6 alkyl group, one or two halogen atoms, C_1-C_6 alkoxy group or CF_1 group;

each $\ensuremath{\mathbb{Q}}$ is independently oxygen atom or sulfur atom;

 Q^1 is oxygen atom or sulfur atom;

 $Z^1 \text{ is } CR^{16}R^{17} \text{ group, oxygen atom, sulfur atom, S(0)} \\$ group, S(0) group or N(C1-C4alky1) group;

each R^{16} is independently hydrogen atom, halogen atom, hydroxyl group, C_1 - C_6 alkoxy group, C_1 - C_6 haloalkyl group, C_1 - C_6 haloalkoxy group, C_2 - C_6 alkylcarbonyloxy group or C_2 - C_6 haloalkylcarbonyloxy group;

each R^{17} is independently hydrogen atom, hydroxyl group or halogen atom;

 R^{10} is $C_1 - C_6$ alkyl group, halogen atom or $C_1 - C_6$ haloalkyl group;

 $$R^{19}$$ and $$R^{20}$$ are independently hydrogen atom, $C_1\!-\!C_6$ alkyl group, or $C_1\!-\!C_6$ haloalkyl group;

 $$Z^{2}$$ is oxygen atom, sulfur atom, NR^{9} group or $CR^{2}R^{10}$ group;

 R^{21} and R^{22} are independently $C_1 - C_6$ alkyl group, $C_1 -$

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 C_6 haloalkyl group, C_3 - C_6 alkenyl group, C_3 - C_6 haloalkenyl group, C_3 - C_6 alkynyl group or C_3 - C_6 haloalkynyl group;

 ${\ensuremath{R^{23}}}$ is hydrogen atom, halogen atom or cyano group;

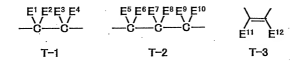
 R^{24} is C_1-C_6 alkylsulfonyl group, C_1-C_6 alkyl group, $C_1-C_6 \ \ \ haloalkyl \ \ \ group, \ \ C_3-C_6 \ \ \ alkenyl \ \ \ group, \ \ C_3-C_6 \ \ \ alkynyl \ \ \ group, \ \ C_1-C_6 \ \ alkoxy \ \ group, \ \ C_1-C_6 \ \ haloalkoxy \ \ group \ \ or \ \ halogen \ \ atom;$

 $R^{25} \mbox{ is } C_1^-C_6 \mbox{ alkyl group, } C_1^-C_6 \mbox{ haloalkyl group, } C_3^-C_6 \mbox{ alkenyl group or } C_3^-C_6 \mbox{ alkynyl group;}$

 R^{26} is $C_1^-C_6$ alkyl group, $C_1^-C_6$ haloalkyl group or phenyl group optionally substituted with $C_1^-C_6$ alkyl, one or two halogen atoms, one or two nitro groups, $C_1^-C_6$ alkoxy group or CF_3 group;

W1 is nitrogen atom or CH group;

T is a group represented by any one of the following general formulas T-1, T-2 and T-3;



(wherein E^1 , E^2 , E^3 , E^4 , E^5 , E^6 , E^7 , E^8 , E^9 , E^{10} , E^{11} and E^{12} are independently hydrogen atom or C_1-C_3 alkyl group);

 R^{27} is $C_1 - C_8$ alkyl group, $C_3 - C_8$ cycloalkyl group,

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C₃-C₈ alkenyl group, C₃-C₈alkynyl group, C₁-C₈ haloalkyl group, C2-C8 alkoxyalkyl group, C2-C8 alkylthioalkyl group, C2-C8 alkylsulfinylalkyl group, C_2 - C_8 alkylsulfonylalkyl group, C_1-C_8 alkylsulfonyl group, phenylsulfonyl group whose phenyl ring may be substituted with at least one substituent selected from the group consisting of halogen atom and C1group, C4-C8 alkoxyalkoxyalkyl group, cycloalkylalkyl group, C6-C8 cycloalkoxyalkyl group, alkenyloxyalkyl group, C4-C8 alkynyloxyalkyl group, C3-C8 haloalkoxyalkyl group, C4-C8 haloalkenyloxyalkyl group, C4-C8 haloalkynyloxyalkyl group, C6-C8 cycloalkylthioalkyl group, C4-C8 alkenylthioalkyl group, C4-C8 alkynylthioalkyl group, C,-C, alkyl group substituted with phenoxy group whose ring is substituted with at least one substituent selected from the group consisting of halogen atom, C1-C3 alkyl group and C,-C, haloalkyl group, benzyloxy group whose ring substituted with at least one substituent selected from the group consisting of halogen atom, C1-C3 alkyl group and C1-C3 haloalkyl group, C4-C8 trialkylsilylalkyl group, cvanoalkvl group, C3-C8 halocycloalkyl group, C2-C0 C5-C8 alkoxyalkenyl haloalkenyl group, group, C₅-C₈ haloalkoxyalkenyl group, C₅-C₈ alkylthioalkenyl group, C₅-C₈ haloalkynyl Cs-Cs alkoxyalkynyl group, group, haloalkoxyalkynyl group, C5-C8 alkylthioalkynyl group, C2-C8 alkylcarbonyl group, benzyl group whose ring is substituted

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with at least one substituent selected from the group consisting of halogen atom, C_1 – C_3 alkyl group and C_1 – C_3 haloalkyl group, $CHR^{34}COR^{28}$ group, $CHR^{34}COOR^{28}$ group, $CHR^{34}P(O)$ (OR^{28})₂ group, $CHR^{34}P(O)$ (OR^{28})₂ group, $CHR^{34}P(O)$ (OR^{28})₃ group, $OR^{28}P^{30}$ group or $OR^{28}P^{30}$ group;

 $R^{28} \mbox{ is } C_1-C_6 \mbox{ alkyl group, } C_2-C_6 \mbox{ alkenyl group, } C_3-C_6$ alkynyl group or tetrahydrofuranyl group;

 $$R^{29}$$ and $$R^{31}$$ are independently hydrogen atom or $C_1\text{--}$ C_4 alkyl group;

 R^{30} and R^{32} are independently $C_1\text{-}C_4$ alkyl group or phenyl group whose ring may be substituted with at least one substituent selected from the group consisting of halogen atom, $C_1\text{-}C_3$ alkyl group and $C_1\text{-}C_3$ haloalkyl group; or,

 $$R^{29}$$ and $$R^{30}$$ together may form $-(CH_2)_5-$, $-(CH_2)_4-$ or $-CH_2CH_2OCH_2CH_2-$, or the ring thus formed may be substituted with at least one substituent selected from the group consisting of C_1-C_3 alkyl group, phenyl group and benzyl group; or,

 $$R^{31}$$ and $$R^{32}$$ may from C_3-C_{ϑ} cycloalkyl group together with the carbon atom to which they are attached;

 $R^{33} \ \mbox{is} \ C_1-C_4 \ \mbox{alkyl group,} \ C_1-C_4 \ \mbox{haloalkyl group or}$ $C_4-C_6 \ \mbox{alkenyl group;}$

 $$R^{34}$$ and $$R^{35}$$ are independently hydrogen atom or $C_1\text{--}$ C_4 alkyl group;

 R^{36} is hydrogen atom, $C_1\text{--}C_6$ alkyl group, $C_3\text{--}C_6$

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alkenyl group or C3-C6 alkynyl group;

 R^{37} is hydrogen atom, $C_1 - C_4 \; alkyl$ group or halogen atom;

 R^{36} is hydrogen atom, C_1 - C_6 alkyl group, C_3 - C_6 cycloalkyl group, C_3 - C_6 alkenyl group, C_3 - C_6 alkoxyalkyl group, C_1 - C_6 haloalkyl group, phenyl group whose ring may be substituted with at least one substituent selected from the group consisting of halogen atom, C_1 - C_4 alkyl group and C_1 - C_4 alkoxy group, $-CH_2CO_2(C_1$ - C_4 alkyl) group or $-CH(CH_3)CO_3(C_1$ - C_4 alkyl) group;

 R^{39} is hydrogen atom, C_1-C_2 alkyl group or $C(0)\,O(C_1-C_4$ alkyl) group;

 $R^{40} \ \ \text{is hydrogen atom,} \ \ C_1-C_6 \ \ \text{alkyl group,} \ \ C_1-C_6$ alkoxy group or NH(C_1-C_6 alkyl) group;

 R^{41} is C_1 - C_6 alkyl group, C_1 - C_6 haloalkyl group, C_1 - C_6 alkoxy group, $NH(C_1$ - C_6 alkyl) group, phenyl group whose ring may be substituted with one substituent selected from the group consisting of R^{42} group, benzyl group and C_2 - C_8 dialkylamino group; and

 $R^{42} \mbox{ is } C_1-C_6 \mbox{ alkyl group, one or two halogen atoms,}$ $C_1-C_6 \mbox{ alkoxy group or } CF_3 \mbox{ group;}$

(3) a compound of the formula (II):

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or nipilacrofen,

wherein R43 is C1-C4 alkyl group;

 R^{44} is C_1-C_4 alkyl group, C_1-C_4 alkylthio group, C_1-C_4 alkoxy group, C_1-C_4 haloalkyl group, C_1-C_4 haloalkylthio group or C_1-C_4 haloalkoxy group;

 R^{43} and R^{44} together may form $-\left(CH_{2}\right)_{3}-$ or $-\left(CH_{2}\right)_{4}-$; R^{45} is hydrogen atom or halogen atom;

 R^{46} is hydrogen atom or C_1-C_4 alkyl group;

 R^{47} is hydrogen atom, nitro group, cyano group, -COOR⁴⁹ group, -C(=X)NR⁵⁰R⁵¹ group or -C(=X²)R⁵² group;

 R^{16} is hydrogen atom, halogen atom, cyano group, C_1 - C_4 alkyl group optionally substituted with at least one substituent selected from the group consisting of halogen atom and hydroxyl group, C_1 - C_4 alkoxy group, phenyl group optionally substituted with at least one substituent selected from the group consisting of halogen atom, nitro group, cyano group, C_1 - C_4 alkyl group, C_1 - C_4 alkoxy group and halo- C_1 - C_4 alkyl group, pyrrolyl group, C_2 - C_8 alkyl group, C_3 - C_8 alkenyl group, C_3 - C_8 alkyl group, C_3 - C_8 alkenyl group, C_3 - C_8 alkenyl group, C_3 - C_8 alkenyl group, C_3 - C_8 alkyl group and C_3 - C_8 alkoxy group into which at least one oxygen atom is inserted, or any one of groups represented by the following formulas:

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wherein $R^{49},\ R^{50}$ and R^{52} are, the same or different, hydrogen atom or $C_1{-}C_4$ alkyl group;

 $$R^{50}$$ and $$R^{51}$$ may form saturated alicyclic 5 or 6 membered ring together with the nitrogen atom to which they are attached;

 $$\rm R^{52}$$ is hydrogen atom, $\rm C_1\text{--}C_4$ alkyl group or $\rm C_1\text{--}C_4$ alkyl group substituted with at least one halogen atom;

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 R^{53} is hydrogen atom, C_1 - C_4 alkyl group optionally substituted with at least one halogen atom, C_2 - C_6 alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl group optionally substituted with at least one halogen atom, phenyl group optionally substituted with at least one halogen atom, C_3 - C_6 cycloalkyl group, cyanomethyl group, or R^{63} CO- group;

 R^{34} is hydrogen atom, C_1 - C_6 alkyl group optionally substituted with at least one halogen atom, C_2 - C_6 alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl group optionally substituted with at least one halogen atom, phenyl group optionally substituted with halogen atom, C_3 - C_6 cycloalkyl group, cyanomethyl group, C_1 - C_4 alkoxy- C_1 - C_6 alkyl group, di- C_1 - C_4 alkylamino- C_1 - C_4 alkyl group, tetrahydrofurfurylmethyl group, C_3 - C_6 alkynyloxy- C_1 - C_4 alkyl group, benzyl whose ring may be substituted with substituent selected from the group consisting of halogen atom, nitro group, cyano group, C_1 - C_4 alkyl group, C_1 - C_4 alkoxy group and halo- C_1 - C_4 alkyl group, $-C_1$ - C_4 alkoxy group and halo- C_1 - C_4 alkyl group, $-C_1$ - C_4 alkoxy group and halo- C_1 - C_4 alkyl group, $-C_1$ - C_4 alkoxy group and halo- C_1 - C_4 alkyl group, $-C_1$ - C_4 alkoxy group, $-C_1$ - C_4 group, $-C_1$ - C_4 alkyl group, $-C_1$ - C_4 alkoxy group, $-C_1$ - C_4 alkyl group, $-C_1$ - C_4 alkoxy group, $-C_1$ - C_4 alkyl group, $-C_1$ - C_4 alkoxy group, $-C_1$ - C_4 alkyl group, $-C_1$ - C_4 alkoxy group, $-C_1$ - C_4 alkyl group, $-C_1$ - C_4 alkoxy group, $-C_1$ - C_4 alkyl group, $-C_1$ - C_4 -

 R^{53} and R^{54} together with the nitrogen atom to which they are attached may form saturated alicyclic 3, 5 or 6 membered ring or aromatic 5 or 6 membered ring in which a carbon atom may be optionally replaced with oxygen

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atom;

 R^{55} is hydrogen atom, C_1-C_4 alkyl group, C_2-C_6 alkenyl group or C_3-C_6 alkynyl group, or R^{55} and R^{56} together may form -(CH₂)_a-;

 R^{56} and R^{57} are independently C_1 - C_4 alkyl group optionally substituted with at least one halogen atom, C_2 - C_6 alkenyl group optionally substituted with at least one halogen atom, C_3 - C_6 alkynyl optionally substituted with at least one halogen atom or phenyl group optionally substituted with at least one halogen atom, hydrogen atom, C_3 - C_6 cycloalkyl group, -XR⁶⁰ group or -NR⁶¹R⁶² group;

 R^{56} is hydrogen atom, C_1-C_6 alkyl group, C_2-C_6 alkenyl group, C_3-C_6 alkynyl group, C_1-C_4 alkylcarbonyl group, cyano- C_1-C_3 alkyl group, C_1-C_4 alkoxycarbonyl- C_1-C_4 alkyl group, $di-C_1-C_4$ alkoxycarbonyl- C_1-C_4 alkyl group, benzyl group, C_1-C_4 alkoxy- C_1-C_4 alkynyl group, $-(CH_2)_a-R^{75}$ group, $-(CH_2)_a-X^2-R^{72}$ group, $-(CH_2)_a-X^2-(CH_2)_a-X^2-(CH_2)_a-X^2-(CH_3)_a$

 $$R^{59}$$ is hydrogen atom, C_1-C_4 alkyl group, C_2-C_6 alkenyl group, C_3-C_6 alkynyl group, cyano- C_1-C_3 alkyl group, C_1-C_4 alkylcarbonyl- C_1-C_3 alkyl group or phenyl group;

 $$R^{60}$$ is $C_1\text{--}C_4$$ alkyl group optionally substituted with at least one halogen atom;

 R^{61} and R^{62} are, the same or different, hydrogen atom or $C_1\text{--}C_4$ alkyl group;

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 R^{63} is C_1-C_4 alkyl group optionally substituted with at least one halogen atom, C_1-C_4 alkoxy- C_1-C_4 alkyl group, C_1-C_4 alkylthio- C_1-C_4 alkyl group, C_3-C_6 cycloalkyl group, phenyl group whose ring may be substituted with one substituent selected from the group consisting of halogen atom, nitro group, cyano group, C_1-C_4 alkyl group, C_1-C_4 alkoxy group and halo- C_1-C_4 alkyl group, $-NR^{73}R^{74}$ group or $-(C_1+C_2)_{1-1}(O)_{1-1}R^{75}$ group;

 $\label{eq:Relation} R^{64} \quad \text{is} \quad C_1\text{--}C_4 \quad \text{alkoxycarbonyl} \quad \text{group;}$ group;

 $$R^{65}$$ is chloromethyl group, cyanomethyl group, $C_3 C_6$ cycloalkyl group into which at least one oxygen atom may be inserted, or C_1-C_4 alkoxycarbonyl- C_1-C_4 alkyl group;

 R^{66} is hydroxyl group or $-NR^{67}R^{68}$ group;

A is $-NR^{67}R^{68}$ group or $-S(0)_f-R^{69}$ group;

 R^{67} and R^{68} are, the same or different, hydrogen atom or $C_1\text{--}C_4$ alkyl group;

 R^{69} is C_1-C_4 alkyl group or C_1-C_4 haloalkyl group;

 R^{70} is hydrogen atom, hydroxyl group, halogen atom, C_1 - C_4 alkyl group optionally substituted with at least one C_1 - C_4 alkoxy group, C_3 - C_6 cycloalkyl group into which at least one oxygen atom may be inserted, C_3 - C_6 cycloalkyl group optionally substituted with one or two methyl groups, furyl group, thienyl group or -C(=0) R^{71} group;

 R^{71} and R^{72} are, the same or different, $C_1\text{--}C_4$ alkyl

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group or C1-C4 alkoxy group;

 $$R^{73}$$ and $$R^{74}$$ are, the same or different, $C_1\text{--}C_4$ alkyl group or phenyl group;

 R^{75} is C_3-C_6 cycloalkyl into which at least one oxygen atom may be inserted, C_3-C_6 cycloalkyl group optionally substituted with one or two methyl groups, furyl group, thienyl group or -C (=0) R^{71} group;

R76 is C1-C4 alkyl group;

a, b and c is independently 1, 2 or 3;

d is 0 or 1;

e is 2 or 3;

f is 1 or 2; and

 X^2 is oxygen atom or sulfur atom.

52. The method according to claim 1,

additionally comprising the steps of:

introducing into the plant cell, a second gene selected from a gene encoding a protein substantially having protoporphyrinogen oxidase activity, a gene encoding a protein substantially having 5-enolpyruvylshikamate-3-phosphate synthase activity and a gene encoding a protein substantially having glyphosate oxidoreductase activity; and

expressing said second gene.

53. A plant cell having:

a gene encoding a protein having the following

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characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin; and
- at least one altered form of an enzymatic activity which gives a resistance to a weed control compound in an amount inhibiting a naturally occurring form of said enzymatic activity, wherein said altered form of an enzymatic activity is a form of enzymatic activity selected from a protoporphyrinogen oxidase activity, 5-enolpyruvylshikamate-3-phosphate synthase activity and glyphosate oxidoreductase activity.
 - 54. A plant cell having:
- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of
 25 modifying a substance for which said protein has a specific

an altered protoporphyrinogen oxidase activity

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affinity, and

(c) being substantially free from framework regions of variable regions in an immunoglobulin; and

which gives a resistance to a weed control compound in an amount inhibiting a natural occurring protoporphyrinogen oxidase activity.

55. A plant cell having:

a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin; and an altered 5-enolpyruvylshikamate-3-phosphate synthase activity which gives a resistance to a weed control compound in an amount inhibiting a natural occurring 5-enolpyruvylshikamate-3-phosphate synthase
 - 56. A plant cell having:

a gene encoding a protein having the following

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activity.

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characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin; and

an altered glyphosate oxidoreductase activity which gives a resistance to a weed control compound in an amount inhibiting a natural occurring glyphosate oxidoreductase activity.

- 57. The plant cell according to claim 53, wherein said altered form of an enzymatic activity is conferred by a second gene selected from a gene encoding a protein substantially having a protoporphyrinogen oxidase activity, a gene encoding a protein substantially having 5-enolpyruvylshikamate-3-phosphate synthase activity and a gene encoding a protein substantially having glyphosate oxidoreductase activity.
- 58. The plant cell according to claim 57, wherein the gene encoding a protein having the following characteristics (a) to (c):
 - (a) having a specific affinity for a substance

which is concerned with the weed control activity of a weed control compound,

- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin; and

the second gene are introduced into the plant cell in the form in that both of said genes are operably ligated to a promoter and a terminator both of which are functional in said plant cell.

- 59. The plant cell according to claim 57, wherein the protein substantially having a protoporphyrinogen IX oxidase activity is protoporphyrinogen IX oxidase, the protein substantially having a 5-enolpyruvylshikamate-3-phosphate synthase activity is 5-enolpyruvylshikamate-3-phosphate synthase and the protein substantially having glyphosate oxidoreductase activity is glyphosate oxidoreductase.
- 60. The plant cell according to claim 53, wherein the plant cell is derived from dicotyledones or monocotyledones.
- \$ 61. A plant comprising the plant cell of claim \$ 54.
 - 62. A plant comprising the plant cell of claim

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55.

- 63. A plant comprising the plant cell of claim 56.
- 64. A method for protecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound to a growth area of the plant of claim 61.
 - 65. A method for protecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of claim 62.
 - 66. A method for protecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of claim 63.
- 67. A method for selecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound to a growth area of the plant of claim 61 and other plants, and selecting either plant on the basis of difference in growth between the plants.
- 68. A method for selecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-

enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of claim 62 and other plants, and selecting either plant on the basis of difference in growth between the plants.

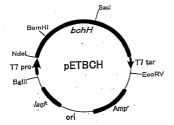
69. A method for selecting a plant which comprises applying a protoporphyrinogen IX oxidase inhibitory-type compound and a compound inhibiting 5-enolpyruvylshikamate-3-phosphate synthase to a growth area of the plant of claim 63 and other plants, and selecting either plant on the basis of difference in growth between the plants.

Abstract of the disclosure:

Weed control compound-resistant plants are produced by introducing a gene encoding a protein having the following characteristics (a) to (c):

- (a) having a specific affinity for a substance which is concerned with the weed control activity of a weed control compound,
- (b) having substantially no capability of modifying a substance for which said protein has a specific affinity, and
- (c) being substantially free from framework regions of variable regions in an immunoglobulin, into a plant cell, and expressing the gene.

- Fig. 1



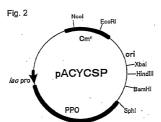


Fig. 3

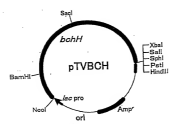
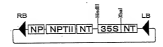


Fig. 4



Fig. 5



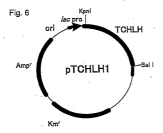


Fig. 7

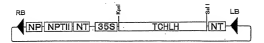


Fig. 8

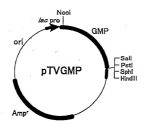


Fig. 10

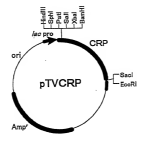




Fig. 12

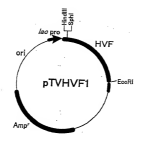


Fig. 13

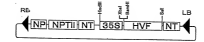


Fig. 14

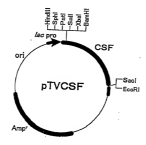
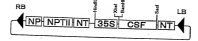
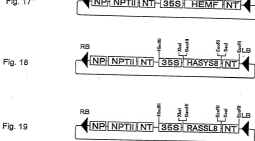
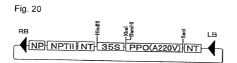
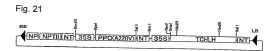


Fig. 15









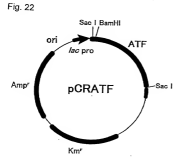


Fig. 23



Fig. 24

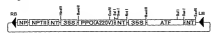


Fig. 25

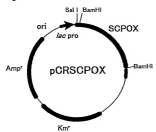


Fig. 26

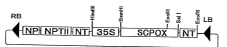


Fig. 27

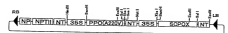


Fig. 28

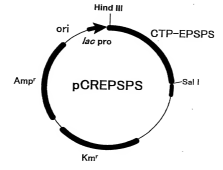


Fig. 29

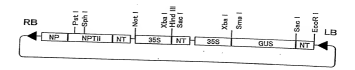
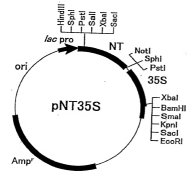


Fig. 30



Fig. 31



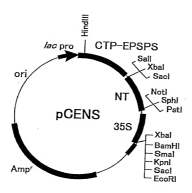


Fig. 33

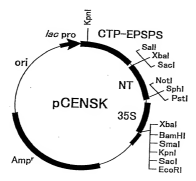


Fig. 34

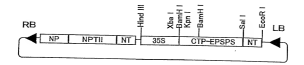


Fig. 35

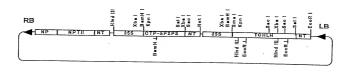


Fig. 36

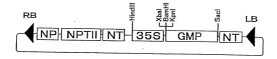


Fig. 37

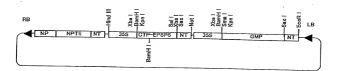


Fig. 38

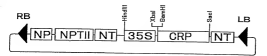


Fig. 39

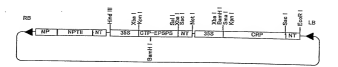


Fig. 40

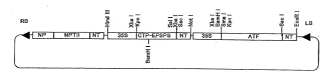


Fig. 41

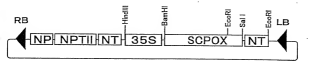
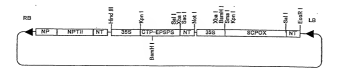


Fig. 42



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ATTORNEY DOCKET NO. 20-4764P

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Page 1 of 2

Telephone: (703) 205-8000 • Facsimile: (703) 205-8050 COMBINED DECLARATION AND POWER OF ATTORNEY FOR PATENT AND DESIGN APPLICATIONS

As a below named inventor, I hereby declare that: my residence, post office address and citizenship are as stated next to my name; that I verily believe that I am the original, first and sole inventor (if only one inventor is named below) or an original, first and joint inventor (if plural inventors are named below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

sert Title: ->	METHOD FOR GIVIN	IG RESISTANCE TO	WEED CONTROL COM	POUNDS TO					
	PLANTS								
I in Appropriate formation —	the specification of which is attached	hereto. If not attached hereto,							
Use →	the specification was filed on as								
hout	United States Application	ion Number		;					
cification	and amended on		(if app	licable); and/or					
ched:	the specification was file	i on		as PCT					
				; and was					
	amended under PCT Art	icle 19 on		(if applicable)					
	I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations,								
1	§1.56.								
3	I do not know and do not believe the same was ever known or used in the United States of America before my or our invention								
n	thereof, or patented or described in any printed publication in any country before my or our invention thereof or more than one year prior								
		to this application, that the same was not in public use or on sale in the United States of America more than one year prior to this application, that the invention has not been patented or made the subject of an inventor's certificate issued before the date of this							
			an application filed by me or my les						
4			tion, and that no application for pate						
nd.			America prior to this application by r						
9	or assigns, except as follows.								
38			ites Code, §119 (a)-(d) of any foreig						
jezh			foreign application for patent or in	ventor's certificate having a					
in the	filing date before that of the applic			n					
rt Priority	Prior Foreign Application	i(s)		Priority Claimed					
imation: ->	120553/1998	Japan	04/30/1998	_ 🖾 🗅					
ppropriate)	(Number)	(Country)	(Month / Day / Year Filed)	Yes No					
	281127/1998	Japan	10/02/1998						
	(Number)	(Country)	(Month / Day / Year Filed)	Yes No					
	330981/1998	Japan	11/20/1998	X □ □ Ves №					
	(Number)	(Country)	(Month / Day / Year Filed)						
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	I haraby alaim the basefit wadon Ti	do 25 Haited Stotes Code \$110(a)	of any United States provisional an	alication(s) listed below					
ert Provisional	I hereby claim the benefit under Title 35, United States Code, §119(e) of any United States provisional application(s) listed below.								
olication(s): -> any)	(Application Number)			(Filing Date)					
	(Application Number) (Filing Date)								
	All Foreign Applications, if any, for any Patent or Inventor's Certificate Filed More than 12 Months (6 Months for Designs) Prior to the								
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_	Country	Application	Number Date o	Date of Filing (Month / Day / Year)					
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	I hereby claim the benefit under T	itle 35, United States Code, §120	of any United States and/or PCT ap	plication(s) listed below and					
	insofar as the subject matter of each	insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States and/or PCT application in							
	the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, §1.56 which became available between the filling date of								
	the prior application and the nation								
rt Prior U.S. lication(s):	. 09/302357	04/30/199		ding					
ny) →	(Application Number)			(Status — patented, pending, abundoned)					
		· -							
	(Application Number)	(Filing Dat	e) (Status	natented pending abandoned)					

I hereby appoint the following attorneys to prosecute this application and/or an international application based on this application and to transact all business in the Patent and Trademark Office connected therewith and in connection with the resulting patent based on instructions received from the entity who first sent the application papers to the attorneys identified below, unless the inventor(s) or assignee provides said attorneys with a written notice to the contrary:

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Donald J. Daley	(Reg. No. 34,313)	John W. Bailey	(Reg. No. 32,881)
John A. Castellano	(Reg. No. 35,094)		

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I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Full Name of First or Sole Inventor:	GIVEN NAME	FAMILY NAME	INVENTOR'S SIGNATURE		DATE*			
Insert Name of ->	Hiroki NAKAJIMA Thok Nary			10/3/00				
Insert Date This Document is Signed	Residence (City, State & Country)			CITIZENSHIP	CITIZENSHIP			
Insert Residence -> Insert Chizenship	Nishinomiya-shi, Hyogo-ken, Japan			Japan				
	POST OFFICE ADDRESS (Complete Street Address including City, State & Country)							
Insert Post Office -> Address	2-29-301, Higashiyamadai, Nishinomiya-shi, Hyogo-ken, Jap							
Full Name of Second Inventor, if any: see above	GIVEN NAME FAMILY NAME INVENTOR'S SIGNATURE				DATE*			
	Akitsu	NAGASAWA	Stitui Vegres		10/3/00			
	Residence (City, State & Co	ountry)	The car To get #	CITIZENSHIP				
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	POST OFFICE ADDRESS (C	omplete Street Address including City, St	ate & Country)					
	5-14-8, Sumiyoshiyamate, Higashinada-ku, Kobe-shi, Hyogo-ken, Japan							
Full Name of Third Inventor, if any see above	GIVEN NAME	FAMILY NAME	INVENTOR'S SIGNATURE		DATE*			
	1							
	Residence (City, State & C	ountry)		CITIZENSHIP				
	POST OFFICE ADDRESS (Complete Street Address including City, State & Country)							
Full Name of Fourth Inventor, if any	GIVEN NAME FAMILY NAME INVENTOR'S SIGNATURE		INVENTOR'S SIGNATURE		DATE*			
see above			<u> </u>					
	Residence (City, State & Country)			CITIZENSHIP				
	POST OFFICE ACORESS (Complete Street Address including City, State & Country)							
	ļ							
	GIVEN NAME FAMILY NAME INVENTOR'S SIGNATURE				I DATE*			
Full Name of Fifth Inversor, if any see above	OTEN WANTE TAINET WANTE.							
22	Residence (City, State & C	Country)		CITIZENSHIP				
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Page 2 of 2 (Revised 11-98)	* DATE OF SIGNATURE							